## AUDREY D. GODDARD, Ph.D.

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1993-present

## PROFESSIONAL EXPERIENCE

Genentech, Inc. South San Francisco, CA

Senior Clinical Scientist Experimental Medicine / BloOncology, Medical Affairs

## Responsibilities:

- Acquisition of clinical samples from Genentech's clinical trials for translational research
- Translational research using clinical specimen and data for drug development and
- Member of Development Science Review Committee, Diagnostic Oversight Team, 21 CFR Part 11 Subteam

- Ethical and legal implications of experiments with clinical specimens and data
- Application of pharmacogenomics in clinical trials

Head of the DNA Sequencing Laboratory, Molecular Binlingy Department, Research

- Management of a laboratory of up to nineteen –including postdoctoral fellow, associate scientist, senior research associate and research assistants/associate levels
- Management of a \$750K budget
- DNA sequencing core facility supporting a 350+ person research facility.
- DNA sequencing for high throughput gene discovery, ESTs, cDNAs, and constructs
- Genomic sequence analysis and gone identification
- DNA sequence and primary protein analysis

### Research:

- Chromosomal localization of novel genes
- Identification and characterization of genes contributing to the oncogenic process
- Identification and characterization of genes contributing to inflammatory diseases
- Design and development of schemes for high throughput genomic DNA sequence analysis
- Candidate gene prediction and evaluation

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Head of the DNA Sequencing Laboratory, Molecular Biology Department, Research

DNA sequencing core facility supporting a 350+ person research facility

- Assumed responsibility for a pre-existing team of five technicians and expanded the group into fifteen, introducing a level of middle management and additional areas of research
- Perticipated in the development of the basic plan for high throughput secreted protein discovery program - sequencing strategies, data analysis and tracking, database design
- High throughput EST and cDNA sequencing for new gene Identification.
- Design end implementation of analysis tools required for high throughput gene identification.
- Chromosomal localization of genes encoding novel secreted proteins.

- Genomic sequence scanning for new gene discovery.
- Development of signal peptide selection methods.
- Evaluation of candidate disease genes. Growth hormone receptor gene SNPs in children with Idiopathic short stature

Imperial Cancer Research Fund London, UK with Dr. Ellen Solomon 1989-1992

- Cloning and characterization of the genes fused at the acute promyelocytic leukemia translocation breakpoints on chromosomes 17 and 15.
- Prepared a successfully funded European Union multi-center grant application

McMaster University Hamilton, Ontario, Canada with Dr. G. D. Sweeney 1983

## 5/83 – 8/83: NSERC Summer Student

In vitro metabolism of β-naphthoflavone in C57BI/6J and DBA mice

### EDUCATION

"Phenotypic and genotypic effects of mutations in the human retinoblastoma gene." Supervisor: Dr. R. A. Phillips

University of Toronto Toronto, Ontario, Canada. Department of Medical Biophysics.

1989

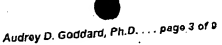
Honours B.Sc

"The In vitro metabolism of the cytochrome P-448 inducer β-naphthoflavone In C57BL/6J mice."

Supervisor: Dr. G. D. Sweeney

McMaster University. Hamilton, Ontario, Canada. Department of Biochemistry

1983



## ACADEMIC AWARDS

Imperial Cancer Research Fund Postdoctoral Fellowship Medical Research Council Studentship NSERC Undergraduate Summer Research Award NSERC Undergraduate Summer Research Award Society of Chemical Industry! Merit Award (Hons. Biochem.) Dr. Harry Lyman Hooker Schnlarship J.L.W. Gill Scholarship Business and Professional Women's Club Scholarship	1989-1992 1983-1988 1983 1983 1981-1983 1981-1982 1980-1981 1979-1980
Business and Processions Scholarship Wyerhauser Foundation Scholarship	

## INVITED PRESENTATIONS

Genentech's gene discovery pipeline: High throughput identification, cloning and characterization of novel genes. Functional Genomics: From Genome to Furirtion, Litchfield Park, AZ., USA. October 2000

High throughput identification, cloning and characterization of novel genes. G2K:Back to Science, Advances in Genome Biology and Technology I. Marco Island, FL, USA. February

Quality control in DNA Sequencing: The use of Phred end Phrep. Bay Area Sequencing 2000 Users Meeting, Berkeley, CA, USA. April 1999

High throughput secreted protein identification and cloning. Tenth International Genome Sequencing and Analysis Conference, Miaml, FL, USA. September 1998

The evolution of DNA sequencing: The Genentech perspective. Bay Area Sequencing Users Mooting, Berkeley, CA. USA. May 1998

Partial Growth Hormone Insensitivity: The role of GH-receptor mutations in Idiopathic Short Stature. Tenth Annual National Cooperative Growth Study Investigators Meeting, San Francisco, CA, USA. October, 1996

Growth hormone (GH) receptor defects are present in selected children with non-GH-deficient short stature: A molecular/basis for partial GH-insensitivity. 76th Annual Meeting of The Endocrine Society, Anahelm, CA, USA. June 1994

A previously uncharacterized gene, myl, is fused to the retinoic acid receptor alpha gene in scute promyelocytic leukemia. XV International Association for Comparative Research on Leukemla and Related Disease, Padua, Italy. October 1991

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#### PATENTS

Godderd A, Godowski PJ, Gurney AL. NL2 Tie ligand homologue polypeptide. Patent Number: 6,455,496. Date of Patent: Sept. 24, 2002.

Goddard A, Godowski PJ and Gumey AL. NL3 Tie ligand homologue nucleic acids. Patent Number: 6,426,218. Date of Patent: July 30, 2002.

Godowski P. Gurney A. Hillan KJ, Botstein D. Goddard A, Roy M, Ferrara N, Tumas D. Schwall R. NL4 Tie ligand homologue nucleic acid. Patent Number: 5,4137,770. Date of

Ashkenazi A. Fong S, Goddard A, Gurney AL, Napler MA, Tumas D, Wood WI. Nurleic acid encoding A-33 related antigen poly peptides. Patent Number: 6,410,708. Date of Patent::

Botstein DA, Cohen RL. Goddard AD, Gurney AL, Hillan KJ, Lawrence DA. Levine AJ, Pennica D, Roy MA and Wood WI. WISP polypeptides and nucleic acids encoding same. Patent Number: 6,387,657. Date of Patent: May 14, 2002.

Goddard A. Gorlowski PJ and Gurney AL. Tie ligands. Patent Number: 6,372,491. Date of

Godowski PJ, Gurney AL, Goddard A and Hillan K. TIE ligand homologue antibody. Patent Number: 6,350,450. Date of Patent: Feb. 26, 2002.

Fong S, Ferrara N. Goddard A, Godowski PJ. Gurney AL, Hillan K and Williams PM. Tie receptor tyrosine kinase Ilgand homologues. Patent Number: 6,348,351. Date of Patent:

Goddard A, Godowski PJ and Gurney AL. Ligand homologues. Patent Number: 6,348,350. Date of Patent: Feb. 19, 2002.

Attie KM, Carlsson LMS, Gesundheit N and Goddard A. Treatment of partial growth hormone Insensitivity syndrome. Patent Number: 6.207,640. Date of Patent: March 27. 2001.

Fong S, Ferrara N, Goddard A, Godowski PJ, Gurney AL, Hillan K and Williams PM. Nucleic acids encoding NL-3. Patent Number: 6,074,873. Date of Patent: June 13, 2000

Attie K, Carlsson LMS, Gesunheit N and Goddard A. Treatment of partial growth hormone Insensitivity syndrome. Patent Number: 5,824,642. Date of Patent: October 20, 1998

Attle K, Carlsson LMS. Gesunheit N and Goddard A. Treatment of partial growth hormone Insensitivity syndrome. Patent Number: 5,646,113. Date of Patent: July 8, 1997

Multiple additional provisional applications filed

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Sechasayee D. Dowd P. Gu Q, Erickson S. Goddard AD Comparetive sequence analysis of PUBLICATIONS the HER2 locus in mouse and man. Manuscript in preparation.

Abuzzahab MJ, Goddard A, Grigorescu F, Lautier C, Smith RJ and Chernausek SD. Human IGF-1 receptor mutations resulting in pre- and post-natal growth retardation. Manuscript in

Aggarwal S, XIe, M-H, Foster J, Frantz G, Stinson J. Corpuz RT, Simmons L, Hillan K, Yansura DG, Vandlen RL, Goddard AD and Gumey AL. FHFR, a novel receptor for the fibroblast growth factors. Manuscript submitted.

Adams SH, Chui C, Schilbach SL, Yu XX, Goddard AD, Grimaldi JC, Lee J, Dowd P, Colman S., Lewin DA. (2001) BFIT, a unique acyl-CoA thioesterase induced in thermogenic brown adipose tissue: Cloning, organization of the human gene, and assessment of a potential link to obesity. Biochemical Journal 360: 135-142.

Lee J. Ho WH. Maruoka M. Corpuz RT. Baldwin DT. Foster JS. Goddard AD. Yansura DG. Vandien RL. Wood WI. Gurney AL. (2001) IL-17E, a novel proinflammatory ligand for the IL-17 receptor homolog IL-17Rh1. Journal of Biological Chemistry 278(2): 1660-1664.

Xie M-H, Aggarwal S, Ho W-H, Foster J, Zhang Z, Stinson J, Wood WI, Goddard AD and Gurney AL. (2000) Interlaukin (IL)-22, a novel human cytokine that signals through the interferon-receptor related proteins CRF2-4 and IL-22R. Journal of Biological Chemistry 275:

Weiss GA, Watanabe CK, Zhong A. Goddard A and Sldhu SS. (2000) Repid mapping of protein functional epitopes by combinatorial alanine scanning. Proc. Natl. Acad. Sci. USA 97:

Guo S, Yamaguchi Y, Schilbach S, Wade T.; Lee J, Goddard A, French D, Handa H. Rosenthal A. (2000) A regulator of transcriptional elongation controls vertebrate neuronal

Yan M, Wang L-C. Hymowitz SG, Schlibach S, Lee J, Goddard A, de Vos AM, Gao WQ, Dixit VM. (2000) Two-amino acid molecular switch in an epithelial morphogen that regulates binding to two distinct receptors. Science 290: 523-527.

Sehl PD, Tai JTN, Hillan KJ, Brown LA, Goddard A, Yang R, Jin H and Lowe DG. (2000) Application of cDNA microarrays in determining molecular phenotype in cardiac growth, development, and response to injury. Circulation 101: 1990-1999.

Guo S. Brush J. Teraoka H. Goddard A. Wilson SW, Mullins MC and Rosenthal A. (1999) Development of noradrenergic neurons in the zebrafish hindbrain requires BMP, FGF3, and the nomeodomain protein soulless/Phox2A. Neuron 24: 555-566.

Stone D, Murone, M, Luoh, S. Ye W, Armanini P, Gurney A, Phillips HS, Brush, J, Goddard A, de Sauvage FJ and Rosenthal A. (1999) Characterization of the human suppressor of fused; a negative regulator of the zinc-finger transcription factor Gli. J. Cell Sci. 112: 4437-

XIR M-H, Halcomb I, Deuel B, Dowd P, Huang A, Vagta A, Foster J, Llang J, Brush J, Gu Q, Hillan K, Goddard A and Gumey, A.L. (1999) FGF-19, a novel fibroblast growth factor with unique specificity for FGFR4. Cytokine 11: 729-735.

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Ridgway JBB, Ng E, Kern JA, Lee J, Brush J, Goddard A and Carter P. (1999) Identification of a human anti-C055 single-chain Fv by subtractive panning of a phage library using tumor and nontumor cell lines. Cancer Research 59: 2718-2723.

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Pennica D, Swanson TA. Welsh JW, Roy MA, Lawrence DA, Lee J, Brush J, Taneyhill LA, Deuel B, Lew M, Watanabe C, Cohen RL, Melhem MF, Finley GG, Quirke P. Goddard AD, Hillan KJ, Gurney AL, Botstein D and Levine AJ. (1988) WISP genes are members of the connective tissue growth factor family that are up-regulated in wnt-1-transformed cells and aberrantly expressed in human colon tumors. *Proc. Natl. Acad. Sci. USA.* 95(25): 14717-14722.

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Xie J. Murone M, Luoh SM, Ryan A, Gu Q, Zhang C, Bonifas JM, Lam CW, Hynes M, Goddard A, Rosenthal A, Epstein EH Jr. and de Sauvage FJ. (1998) Activating Smoothened mutations In sporadic basal-cell carcinoma. *Nature*. 391(6662): 90-92.

Marsters SA, Sheridan JP, Pitti RM, Huang A, Skubatch M, Baldwin D, Yuan J, Gumey A, Goddard AD, Goddwski P and Ashkenazl A. (1997) A novel receptor for Apo2L/TRAIL contains a truncated death domain. Current Biology. 7(12): 1003-1006.

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Marsters SA, Sheridan JP, Donahue CJ, Plttl RM, Gray CL, Goddard AD, Bauer KD and Ashkenazi A. (1996) Apo-3, a new member of the tumor necrosis factor receptor family, contains a death domain and activates apoptosis and NF-kappa β. Current Biology 8(12): 1669-76

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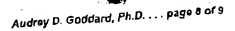
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Molecular cloning of a ligand for the EPH-related receptor protein-tyrosine kinase Htk. Proc.
Natl. Acad. Sci. USA 92: 1866-1870.

Huang X, Yuang J, Goddard A, Foulis A, James RF, Lemmark A, Pujol-Borrell R, Rabinovitch A, Somnza N and Stewart TA. (1995) Interferon expression in the pancreases of patients with type I diabetes. *Diabetes* 44: 658-664.

Goddard AD, Yuan JQ, Fairbairn L, Dexter M, Borrow J, Kozak C and Solomon E. (1995) Cloning of the murine homolog of the leukemia-associated PML gene. *Mammalian Genome* 8: 732-737.



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Kuo SS, Moran P, Gripp J, Armanini M, Phillips HS, Goddard A and Caras IW. (1994) Identification and characterization of Batk, a predominently brain-specific non-receptor protein tyrosine kinase related to Csk. J. Neurosci. Res. 38: 705-715.

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Foulkes W, Goddard A, and Patel K. (1991) Retinoblastoma linked with Seascale [letter]. British Med. J. 302: 409.

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HellerEhrman

ESEARCH/

SIMULTANEOUS AMPLIFICATION AND DETECTION OF SPECIFIC DNA SEQUENCES

To be the Midwide Russell Higuchi\*, Gavin Dollinger¹, P. Scan Walsh and Robert Criffith

Russell Higuchi\*, Gavin Dollinger¹, P. Scan Walsh and Robert Criffith

Russell Higuchi\*, Gavin Dollinger¹, CA 96508, 'Chiron Corporation, 1400 53rd 3tr, Enterpyille, CA

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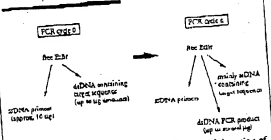
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These downstream processing steps would be climinated if specific are phiseation and detection of amplified DNA took place simultaneously within an unopened reaction vessel. Assays in which such different processes take place without the need to separate reaction components have been termed "homogeneous". No Truly homogeneous PCK assay has been demonstrated to date, although progress towards this end has been reported. Chebab, et al. 12, developed a PCR product detection scheme using

Augrescent primers that resulted in a Augrescent PCR product. Allele-specific primers, each with different fluoresear tage, were used to indicate the xenotype of the DNA. However, the unincorporated primers must still be result. Recently, Holland, et al. 3, developed an assay in which the endogenous 5' exonuclease assay of Taq DNA polymerase was exploited to cleave a labeled oligonucleotide probe. The probe would only cleave if PCR amplification had produced its complementary sequence. In order to detect the cleavage products, however, a subse-

quent process is again needed. We have developed a truly homogeneous assay for PCR who PCR product detection based upon the greatly increased fluorescence that othidium bromide and other DNA binding dyc: exhibit when they are bound to de-DNA 14-18. As outlined in Figure 1, a protocypic PCR



PROBLE & Principle of simultaneous amplification and detection of PCR product. The components of a PCR exemining ERBY that are Proposed to the PCR exemining ERBY that are Proposed to the listed—ERBY itself, ERBY bound to either sidned about the ERBY is dr. DNA. There is a large fluorescence cohencement when ERBY is doubt to DNA and building is growly enhanced when INA is doubt-stranded. After sufficient (a) cycles of PCR, the net increase in dr. DNA results in additional ERBy binding, and a set increase in total fluorescence. increase in total Autorescence.

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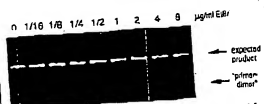
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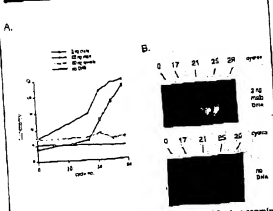
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FROME 1 Gel electrophorets of PCR amplification products of the human, nuclear gene, HLA DQa, made in the presence of increasing amounts of EtBr (up to 5 ug/ml). The protecte of EtBr has no obvious effect on the yield or specificity of amplifications.



PCERT 3 (A) Fluorescence measurements from PCEs that contain 0.5 µg/ml Echr and that are specific for V-chromosome repeat 0.5 µg/ml Echr and that are specific for V-chromosome repeat equances. Five replicate PCEs were begun containing each of the Equation of the PCEs for each DNA was removed from thermosphing and its PCEs for each DNA was removed from thermosphing and its PCEs for each DNA was removed from thermosphing and its PCEs for each DNA was removed from thermosphing and its PCEs for each DNA was removed from Exploration (B) PCEs there (0.6 ml Eppendorf-style, polyprovided in the polyprovided

begins with primers that are single-manded DNA (ss-DNA), dNTPs, and DNA polymersse. An amount of dsDNA containing the target sequence (target fina) is also typically present. This amount can vary, depending on the application, from single-cell amounts of DNA<sup>17</sup> to nurrograms per PCR. It Ethr is present, the reagents that will fluoresce, in order of increasing fluorescence, are free Ethr welf, and Ethr bound to the single stranded DNA primers and to the double-stranded target DNA (by its intercalation bereaen the stacked bases of the UNA double-hells). After the first denaturation cycle, target INA will be largely single-stranded. After a PCR is completed, the most significant change is the increase in the amount of dsDNA (the PCR product itself) of up to several micrograms. Formerly tree EtBr is bound to the additional dsDNA, resulting in an increase in fluorescence. There is also some decrease in the amount of 13DNA primer, but because the binding of EtBr to seDNA is much less than to dsDNA, the effect of this change on the total fluorescence of the sample is small. The fluorescence increase can be measured by directing excitation illumination through the walls of the amplification vessel

before and after, or even continuously during, therinoryding.

PCK in the presence of Ernr. In order to assess the RESULTS affect of Libr in PCR, amplifications of the human HIA antect of Ethr in Puk, ampuneations of the human Ph. A DQa gene's were performed with the dye present at concentrations from 0.06 to 8.0 µg/ml (a typical concentration of Ethr used in staining of nucleic acids following training of the property of the public of the pu gel electrophoresis is 0.5 µg/ml). As shown in Figure 2. gel electrophorais revealed little or no difference in the yield or quality of the amplification product whether Lthr was absent of present at any of these concentrations, indicating that Ethr does not whibit PCR.

Detection of human Y-chromosome specific tequences. Sequence-sponic, fluorescence enhancement of EiBr as a result of PCR was demonstrated in a series of amplifications containing 0.5 ug/ml EtBr and primere specific to repeat DNA sequences found on the human yearing to repeat DNA sequences found on the human yearoncomes. These PCRs initially contained either 60 ng male, 60 ng femalo, 2 ng male human or no DNA. Five replicate PCRs were begun for each DNA. After 0, 17, 21, 24 and 29 cycles of thermocycling, a PCR for each DNA was removed from the thermocycler, and its fluorescence measured in a spectrofluorometer and plotted vs. amplification cycle number (Fig. 3A). The shape of this curve reflects the fact that by the time an interase in the detected, the increase in DNA is becoming linear and not constant and such as the second of becoming linear and not exponential with cycle number. As shown, the fluorescence increased about three-fold over the background Muorescence for the PURS containing human male DNA, but did not significantly increase for negative control PCRs, which contained nither no DNA or human female DNA. The more make HNA present to begin with—60 ng versus 2 ng—the fewer cycles were needed to give a detectable increase in fluorescence. Gel electrophoresis on the products of these amplifications showed that DNA fragments of the care pected size were made in the male DNA containing reactions and that little DNA synthesis took place in the

In addition, the increase in fluorescence was visualized control samples. by simply laying the completed, unopened PCRs on a UV nanilluminator and photographing them through a red filter. This is shown in figure 3B for the reactions that

began with I ng male DNA and those with no DNA.

Detection of specific alleles of the human β-slobin gene. In order to demonstrate that this approach has adequate spacificity to allow genetic screening, a detection of the sickle-cell anemia mutation was pertormed. Figure 4 shows the fluorescence from completed amplifications conceining EtBr (0.5 mg/ml) 23 detected by photography of the reaction tubes on 2 UV cransilluminator. These Teactions were performed using primers specific for elther the wild-type or sickle-cell mutation of the human-by placing the sickle-mutation six at the terminal structure of one primer. By using an appropriate primer annealing temperature, primer excession—and thus among the second of the sec annealing temperature, primer extension—and thus an anneaus an take place only if the 9' nucleonde of the primer is complementary to the 8-globin allele present

Each pair of amplifications shown in Figure 4 consists of a reacoun with either the wild-type allele specific (left rube) or sickle-allele specific (right tube) primers. Three different DNAs were typed: DNA from a homorygous wild type β-globin individual (AA): from a heterozygous eickle β-globin individual (AS); and from a homozygous eickle β-globin individual (SS). Each UNA (51) ng genomic DNA to start each PCR) was analyzed to triplicate (8 pairs

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of rescuent such). The DNA type was reflected in the or relative fluorescence intensities in each pair of completed amplifications. There was a significant increase in fluoresampuncations. There was a significant increase in muores-cance only where a B-globin allele DNA matched the prince see When measured on a spectrofluorometer primer see when measured on a spectromorometer (data not shown), this fluorescence was about three times that present in a PLR where both B-globin alloles were that present in a where total p-grown alloles were shown established that this increase in fluorescence was due to the synthesis of nearly a microgram of a DNA fragment of the expected size for P-globin. There was little synthesis of deDNA in reactions in which the allele-

specific primer was mismatched to both alleless
Continuous monitoring of a PCR. Using a fiber optic device. It is possible to direct excitation illumination from a specifofluorometer to a PCR undergoing thermocycling and to return its fluorescence to the spectrofluorometer. The Augrescence readout of such an arrangement, directed at an EiBr-containing amplification of Y-chromosome specific sequences from 25 ag of human male DNA, is shown in Figure 5. The readout from a control PCR with no target DNA is also shown. Thirty cycles of POR er 0, cach fluo-success f while

were monitored for each. The Suorescence trace as a function of time clearly shows the effect of the thermocycling. Fluorescence intensity rises and falls inversely with temperature. The fluorescence intensity is minimum at the denaturation temperature (94°C) and maximum at the annealing/extension temperature (50°C). In the negative-control PCR, these Ausrescence maxima and minima do not change signifiand number of the control of the con the continuous illumination of the comple.

se ing la suberior fold in the second fluor fluo in the PCK containing maic DNA, the fluorescence maxima at the annealing/extension temperature begin to increase at about 4000 seconds of thormocycling, and continue to increase with time, indicating that de DNA is being produced at a detectable level. Note that the Buoreseence minima at the denaturation temperature do not agnificantly increase, presumably because at this temperature there is no deDNA for ECBr to bind. Thus the course of the amplification is followed by tracking the fluoresective increase at the annealing temperature. Analysis of the products of these two amplifications by gel electropho-resis showed a DNA fragment of the expected size for the male DNA containing nample and no detectable DNA synthesis for the control sample.

Downstream processes such as hybridization to a sc-**DISCUSSION** quence-specific probe can enhance the specificity of DNA disection by FCR. The elimination of these processes means that the specificity of this homogeneous assay depends solely on that of PCR. In the case of sickle-cell disease, we have shown diat PCR along her sufficient DNA

for a significant solely on that of FCR. In the seasificant DNA number of diesse, we have shown that PCR alone has sufficient DNA particles sequence specificity to permit generic acreening. Using sequence specificity to permit generic acreening. Using appropriate amplification conditions, there is little non-interest appropriate target allele.

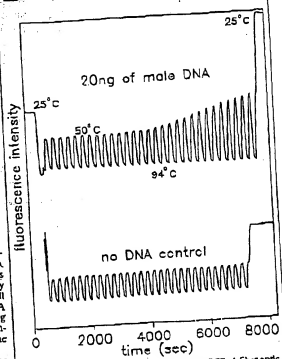
of the specificity required to detect pathogens can be not some or less than that required to do generic acreening, insufficiently the amount of other DNA that must be taken with the teleft of the amount of other DNA that must be taken with the Three states amount of other DNA that must be taken with the teleft of the amount of other DNA that must be taken with the Three states amount of other DNA that must be taken with the teleft of a few copies sygonic per thousands of host cells. Compared with genetic sygonic per thousands of host cells. Compared with genetic specificity and the input of more total to the particular specificity and the input of more total



Homozygous AA Heterozygous AS

> Homozygous SS

Note: 4 UV photography of PCR cubes containing amplifications using EIBr that are specific to wild-type (A) or stekle (S) silleles of the human B-globin sense. The left of each pair of tubes contains allele specific primers to the wild-type alleles, the right tube primers to the sielle allele. The photograph was taken after 30 primers to the sielle allele. The photograph was taken after 30 primers to the sielle allele. They have sued to begin FCR. Trying are indicated. Fifey up of DNA was used to begin FCR. Trying was done in triplicate (3 pairs of PCRs) for each input DNA.



PROBE 5 Condenses, real-time monitoring of a PCR. A fiber optic was used to carry excitation light to a PCR in progres and also emitted light back to a fluoremeter (see Experimental Protocol), amitted light back to a fluoremeter (see Experimental Protocol). Amplification using human male-DNA specific primers in a PCR starting with 20 ng of human male DNA (top), of in a control starting with 20 ng of human male DNA (top), of in a control starting with 20 ng of human male DNA (top), of in a control starting with 20 ng of human male DNA (top), of in a control starting with 20 ng of human male DNA (top), of in a control starting with 20 ng of human male DNA flow (top), of the control starting and extension). Note in the male DNA PCM, the cycle (time) dependent increase in Eutorescence at the annualing/extension temperature.

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THE MADE STATE DNA—up to microgram amounts—in order to have sufficient numbers of target sequences. This large amount of starting DNA in an amplification significantly increases the background fluoreteence over which any additional Autoresience produced by FCR must be detected. An additional emplication that occurs with marges in low copy number is the formation of the "primer-dimer" arniact. This is the result of the extension of one primer using the other primer as a template. Although this occurs infrequently, once it occurs the extension product is a substrate for PCR amplification, and can compete with dimer product is of course de DNA and thus is a potential source of felse signal in this homogeneous assay.

To increase PCR specificity and reduce the effect of

primer dimet amplification, we are investigating 2 number of approaches, including the use of acsted primer amplifications that take place in a single rube, and the "hot-start", in which nonspecific amplification is reduced by raising the temperature of the reaction before DNA synthesis begins. Preliminary results using these approaches suggest that primer-dimer is effectively roduced and it is possible to detect the increase in Ethr fluorescence in a PCR insulgated by a single HIV genome in a background of 102 cells. With larger numbers of cells, the background fluorescence contributed by genomic DNA becomes problement. To reduce this background, it may be possible to use sequence-specific DNA-binding dyes that can be made to preferentially bind PCR product over that can be made to preferentially bind PCR product over that can be made to preferentially bind PCR product over that can be made to preferentially bind PCR product over that can be made to preferentially bind PCR product over that can be made to preferentially bind PCR product over that can be made to preferentially bind PCR product over the discount of the product of the prod genomic DNA by incorporating the dye-building DNA sequence into the PCR product through a 8 "add-on" to

the aligorudeoride primered. we have shown that the detection of fluorescence generated by an ErBr-containing PCR is straightforward. both once PCR is completed and community during thermocycling. The ease with which automation of specific DNA detection can be accomplished is the most promising aspect of this assay. The fluorescence analysis of completed PCRs is already possible with existing instru-mentation in 96-well format. In this format, the fluorescease in each PCR can be quantizated before, after, and even at selected points curing thermocycling by moving the rack of PCRs to a 96-microwell plate huorescence the rack of PCRs to a 96-microwell plate huorescence

The instrumentation necessary to community monitor readeres. multiple PCRs simultaneously is also simple in principle. A direct extension of the apparatus used here is to have multiple fiberoptics trensmit the excitation light and fluorescent emissions to and from multiple PCBs. The ability to monitor multiple PCRs continuously may allow quantitation of target DNA copy number. Figure 3 shows that the larger the amount of starting target DNA, the sconer during PCR a fluorescence increase is detected. Prelimination of the starting target DNA, the sconer during PCR a fluorescence increase is detected. nary experiments (Higuchi and Dollinger, manuscript in preparation) with continuous monitoring have shown a sensitivity to two-fold differences in initial target DNA

Conversely, if the number of target molecules is concentration. known as u can be in genetic screening concinuous monitoring may provide a means of detecting false posttive and false negative results. With a known number of Erget molecules, 2 true positive would exhibit detectable fluoreseence by a predictable number of cycles of PCR. Increases in fluorescence detected before or after that cycle would indicate potential arufacts. False negative results due to, for example, inhibition of DNA pulymerase, may be detected by including within each PCR an inefficiently amplifying marker. This marker results in a fluorescence increase only after a large number of cy--many more than are necessary to detect a true

positive. If a sample fails to have a fluorescence increase after this many cycles, inhibition may be suspected. Since, in this array, conclusions are drawn based on the presence or absence of fluorescence signal alone, such controls may be important. In any event, before any test based on this principle is ready for the clinic, an assessment of its false positive/talse negative rates will need to be obtained using

a large number of known samples.

In summary, the inclusion in POR of dyes whose flucrescence is enhanced upon binding diDNA makes upon bossible to detoct specific DNA amplification from outside the PCR tube. In the future, instruments based upon this principle may facilitate the more wide present use of PCR in applications that demand the high chroughput of

mapples.

\*\*\*CFERIMENTAL PROTOCOL\*\*\*

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uring an annealing/extension temperature of 50°C. The reaction was covered with mineral oil (2 drops) to prevent evaporation. Thermocycling and filtorescence measurement were trained multimeously. A time-base scan with a 10 second insegration of the multimeously.

vas used and the emission signal was radoed to the excitation signal to control for changes in light-source intensity. Data were lollected using the dm3000f, version 2.6 (SPEX) data system.

Admortedgments We thank Bob Jones for help with the spectrofluormetric was transfer and Heatherbell Fong for editing this manuscript.

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Tumosa, N. and Rahan. L. 1989. Eluccocent ElA sectioning of monoclogal andbodies in cell surface antigent. J. Ismaun. Mech. 116:59-58.



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# Oligonucleotides with Fluorescent Dyes at Opposite Ends Provide a Quenched Probe System Useful for Detecting PCR Product and Nucleic Acid Hybridization

Kenneth J. Livak, Susan J.A. Flood, Jeffrey Marmaro, William Giusti, and Karin Deetz Carling Street

Botkin-Elmer, Applied mosystoms Division, Foster City, California 94404

The 5' nucloase PCR easy detects the accumulation of specific PCR product by hybridization and cleavage of a double-labeled fluorogenic probe during the amplification reaction. The probe is an oligonucleotida with both a reporter fluorescent dye and a quencher dye attached. An increase in reporter fluorescence intensity indicates that the probe has hybridized to the target PCR product and has been cleaved by the 5'-3' nucleolytic activity of Tag DHA polymerase. in this study, probes with the quencher dye attached to an internal nucleotide were compared with probes with the quencher dye at tached to the 3'-and nucleotide. In all cases, the reporter dye was attached to the 5' and. All intact probat showed quenching of the reporter fluorescence. In general, probes with the quencher dye attached to the 3'and nucleotide exhibited a larger signal in the 5' nucleate PCR assay than the internally inhaled probac it is proposed that the larger signal is caused by increased lihelihood of cleavage by Tag DHA polymerasc when the probe is hybridized to a template strand during PCR. Probes with the quencher dye attached to the 3'-and nucleatide also exhibited en increase in reporter fluorescence Intensity when hybridized to a com-, plemontary strand. Thus, oligonucleorldes with reporter and quencher dyes attached at opposite ends can be used as homogeneous hybridiza-

A homogeneous usuay for detecting the meaning etton of specific PCR product that uses a double-tabeled filtorogenic probe was described by Levet al. (1) The array exploits the 5' . 3' nucleolytic activity of Tag INA polymease 2.41 and is diagramed in figure 1. The iluarogenic probe consists of an oligonucleotide will a reporter fluorescent dyr, such as a fluorescein, attached to the 5 end and a quencher dye, such as a rhodamine, attached internally, When the fluorescen is excited by irradiation, lis fluorescent emission will be quenched if the dandantine is closs enough to be excited through the pre-COS OL MUDIESCENCE CHARTY transfer bridized to a template strand, Tag DNA polymerase will cleave the probe because of its inherent 5' -- 3' nucleolytic activity. If the cleavage occurs between the fluorescein and rhodamine dyes, it causes on increase in fluctionervire fluores. cence intensity because the fluorescein is no longer quenched. The increase in flourescein fluorescence intensity indicates that the probespectale PCR product has hear generated. Thus, PBT between a reporter dye and a quencher dye is miteat to the performance of the probe in the 5' nuclease I'CR may.

Quenching is completely dependent on the physical proximity of the two dyes. (4) Because of this, it lies been us. sumed that the quencher dye must be altached near the 5' end. Summaingly, we have found that stisching a rhodamine dye at the 3' end of a probe

PCIt assay, Purthermore, cleavage of this type of probats not required to achieve some reduction in quenching, Oligonucleoudes with a reporter dye on the S' and and a quencher dye on the 3' end exhibit a much higher reporter fluorescence when double-stranded as compared with single-stranded. This should make it possible to use this type of doubladabeled probe for nomogeneous detection of nucleic acid hybridization.

## MATERIALS AND METHOUS

## Ollgonucleotides

Table 1 shows the nucleotide sequence of the oligonucleotides used in this stildy, tanker arm nucleotide (LAN) phosphoramidite was obtained from Glen Research. The standard DNA phosphoramidires, 6-carboxylluorescella (6-PAM) phosphoramidite, d-carboxyretmmelhyirhodamine succlinimityl ester (TAMRA NHS exter), and Phosphalink for attaching a 3'-blocking phrisphate, were obtained from Parkin-Simer, Applied Biosystems Division. Oligonucleotide synthesis was performed using an ABI model 394 FINA synthesizer (Applied Blosystems). Primer and complement oligonucleandes were purifica using Olige Postfication Cartridges (Applied Blosystems). Double-labeled probes were synthesized with G-PAM-labeled phosphoramidite at the 5' and, IAN replacing one of the T's in the sequence, and Phosphalink at the 3' end. Pollowing deprotection and charce precipitation, From : BML

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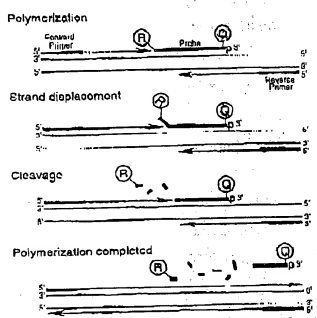


FIGURE 1 Diagram of 5' nuclease away. Seepwise representation of the 5' -- 3' nucleolytic attivity of Tag DNA polymerase acting on a fluorogenic probe during one extension phose of PCR, A Park SA

ms Na-bicarbonate buffer (pll 9.0) at room temperature. Unreacted dye was removed by passage over a ID-10 Scylm. dex column. Finally, the double-labeled probe was purifical by preparative highperformance liquid chromatography (IIPIL) using an Aquapore C. ZZIXA.C. man column with 7-pm particle size. The column was developed with a 24-min linear gradient of 8-20% acctonitelle in U.1 H TEM (triethylamine accesse). Probes are named by designating the sequence from Table 1 and the position of the IAN-TAMRA motery. For example, probe A1-7 has sequence A1 with LAN-TAMRA at nucleotide position 7 from the 5' und.

#### PCR Systems

All PCR amplifications were performed in the Perkin-Elmer GeneAmp PLR System 9600 using 50-ul reactions that contelned 10 mm Tria-HCI (pl) 8.3), 50 mm КСІ, 200 нм фЛТР, 200 нм фСТГ, 200 нм dGTP, 400 per dUTP, 0.5 unit of Amperuse uracil N-elycosylase (Perkin-Limer). gene (nucleotldes 2141-2435 in the sequence of Nakailina-Illiana et al.) Was TAN bus YIV coming guize belilque (Table 1), which are modified slightly from those of du Breuil et al. (10) Actin amplification reactions commined 4 mm MgCl<sub>p</sub>, 20 ng at human genomic BNA, 1020 MM OUE bas, sdorg EA to IA MM 02ch primer. The thermal regimen was SO'C (2 min), 95°C (10 min), 40 cycles of 95°C (20 sec), corc (1 min), and hold at 72°0. A 515-bp segment was amplified from a plasmid that consists of a segment of & DNA (nucleotides 32,220-32,747) inserted in the Smal site of vector pUC119: These reactions contained 5.5 ms MECIZI 1 ng of plusmid DNA, 50 rm P2 or PS probe, 200 is primer F110, and 200 um piliner R119. The thermal regimen Was 50°C (2 min), 95°C (10 min), 25 cydes of 95°C (20 sec), 57°C (1 min), and hold at 72°C.

#### Munrescence Detection

For each amplification reaction, a actual aliquol of a sample was transferred to an Individual well of a white, 96-wall interotiter plate (Perkin-Fimer). Fluorescence was measured on the Perkin-Umer Tag-Man LS-50B System, which consists of a luminescence spectrometer with plate reader astembly, a 483-nm excitation filter, and a 515-nm emission filter. Pacitation was at 488 nm using a 5-nm slit width, limission was measured at 518 am for 6-PAM (the reporter or R value) and SRI am for TAMILA (the guencher of Q value) using a Iti-nm elit width. To determine the increase in reporter embsion that is caused by cleavage of the probe during PCK, three nurmalizations are applied to the raw emissions date. First, emission intensity of a buffer blank Is subtracted for each wavelength, Secand, emission intensity of the reporter is

TABLE 1 Sequences of Oligonucleotides

Name	1уре	Zequence
P119 R119 P2 P2C P5 INC ARP ARP A1 A1 A1 A3 A3C	primer primer probe complement probe primer	ACCEACAGAACTGATCACCACTC ATCTCTTGCTCGAAGTGCAGGGGAAC CATCTCTTGCTCGAAGATGCAGAGATACCACTP ACCCACAGAGACTTGCAACATTCCAACAATACAATACAA

For each oligonucleonide used in this study, the nucleic add sequence is given, written in the " dieserton. There are three tween of clicumucleotides: PCR primer, fluorogenic probe used rom : BML

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A1-2	RACINGROCCCCARCITATCCTCCCTP
A1-7	PARCECCOCCECATOUCATCETCCCTIV
A1-14	MAINGCOMPCCCONOCCENICCTOCTO
A1-10	אישבבביבבבבהישבבי לבהוספמים
A1-22	RATGCCCTCCCCCNTCCCNTCCQCCCNT
A1-26	PANCOUTTOCCOTATERIO TOCTOCTO
	A1-7 A1-14 A1-10 A1-22

Probin	618 nm		582 nm		RQ.	RQ I	ARO
	no temp,	4 temp.	no semp	_ a temp			
	30.0 L 2.1	32,7 ± 1.0	20.2 4 0.0	68.2 - 2.9	067+0.01	20.0 1,02.0	9.18 4 0.00
A1:2		306.1 a 21.4	•	1102 - 63	640 - 0.02	2.58 - 0.17	360 7 C.18
۸1-7	53.0 ± 4.3	306.13 212		i kana	1.0100	4944618	3.18 ( 0.15
A1-14	127.0 + 4.9	433.5 + 18.1	100.7 ± 8.3	93.1462	1.0300	0,10 ي مرزو	
A1-19		400.71 7.7	70.2 + 7.4	78.0 4. 0.0	2.67 a 0.06	21.0 1 00'S	3.12 : 6,16
	224.04 0.4	480.V e 43.6	100.0 ± 4.0	86.2 1 0.6	£.25 ± 0.03	5.0210.11	2,77 ± 0.12
A1-22	224.04 04	49U,8 8 43.0				501 ± 0.05	238 ± 0.08
Δ1-28	160.21 0.9	44.1 1 18.4	93.1 ± 5.4	81.7 ± 3.8	1.12 2 0 02	5,0120.00	320,20.00

ROURE 2 Results of & mucleose reserve uniqueing Fracity probes with TAMEA at different nucle outle positions. As described in Materials and Methods, PCIC simplifications containing the indicated probet were performed, and the fluorescence emission was measured at 514 and 382 nm. Reported values are the average=1 s.p. for six reactions run without added template (no temp.) and six reactions run with templace (4 tump.). The RQ ratio was calculated for each individuel wastion and anciating to live the seborted RO, and HO, August

givided by the emission intensity of the quencher to give an RQ ratio for coch isaction tube. This normalizes for wellto-well veriations in probe connectes. non and fluorescence measurement. Ptnamy, area in calculated by subtracting the KQ value of the no-template control [RQ"] from the RQ value for the compiete reaction including template (RQ').

#### RECURTS

A series of probes with increasing diswires between the unprescent tehonics and rhodemine quenches were tested to investigate the minimum and maximum spacing that would give an acceptable performance in the S' nuclease I'CH assay. These probes hybridize to a target sequence in the human p-acrin gene. Figure 2 shows the results of an exportment in which these probes were included in PCR that amplified a segment of the Barilli galla containing the larget sequence Performance in the 5' auclease PCR away is monitored by the magnitude of ARO, which is a measure of the increase in reporter fluorescenice caused by PCR amplification of the probe turger, Probe A1-2 has a SRQ value that is close to zero, indicating that the probe was not cleaved appreciably tiuring the amplification reaction. This sug-Keals that wills the quanches dye on the secund nucleating from the 5' and, there la insufficient fount for Tay polymerase to cleave efficiently between the reporter and quenches. The other five probes exhillited comparable ARC values that are

clearly different from zero. Thus, all five profies are being cleaved during PCR amphilication resulting in a similar increase in reporter fluorescence. It should be noted that complete digestion of a proba produces a much larger increase in reporter fluorescence than that observed in Figure 2 (data not thown). Thus, even in reactions where amplification occurs, the majority of probe molecules remain uncleaved. It is mainly for this reason that the fluorescence intentity of the quencher dye TAMILA changes lillie with amplification of the target. This is what allows us to use the 582-rim fluorescence. reading as a normalization factor.

The magnitude of RQ dopende mainly on the quenching efficiency inherent in the specific afficience of the probe and the purity of the oligonucleotide. Thus, the larger HQ - values indicate that probes A1-14, A1-19, A1-22, and A1-28 propably have reduced quenching as compared with A1-7. 3th, the degree of quenching it sufficient to detect a highly significant increases in reporter fluorescence when each of these probes

is cleaved during PCR.

To further investigate the ability of TAMKA on the 31 and to quanch G-PAM on the 3' end, three additional pairs of probes were tested in the 5' nuclease PCR ussay. For each pair, one probe has TAMRA etteched to an internal nuclawilde and the other has TAMM attached to the 3' end nucleotide. The results are shown in Table 2. For all three sets, the probe with the 3' quencher exhibits a ARQ value that is considerably higher than for the probe with the internal quencher. The RQ values suggest that differences in quanching are not as great as those abserved with some of the Al probes. These results demonstrate that a quencher dyc on the 3' end of an oligonucleatide can quench efficiently the

The second of th TABLE Z Kesuits of 5' Nuclease Assay Comparing Probin with TAMRA Attached to an Internal or 3'-terminal Nucleated

318 nu		ntii	582 nm			501	AKU
			na temp.	+ temp.	NQ	RQ'	
Probe	no temp.	+ tewl-			0.47 ± 0.62.	0,73 = 0.0%	0.26 & 0.03
	\$4.6 1 3.2	84.8 Z 3.7	110.2 = 0.4	175.6 ± 2.5 90.2 ± 3.8	0.86 T 0.05	2.62 = 0.05	$1.76 \pm 0.09$
A3-6 A3-24	72.1 + 2.9	236.5 ± 11.1	RA.2 ± 4.0	•	0.79 T OTS	3.19 × 0.16	2.40 ± 0.1
127	62.8 3. 4.4	384.U ± 34.1	105.1 = 6.4	120.4 = 10.2 118.7 = 4.8	0.81 ± 0.01	4.68 ± 0.10	3.68 = 0.1
12-27	113.4 = 6.6	556.4 4 14.1	.190.7 = 8.5		0,89 = 0,08	2.55 5. 0.06	$1.60 \pm 0.0$
•	77,3 = 6.5	244.4 a 15.9	PA.7 1. 4.3	95.8 = 0.7 94.7 = 6.3	(L63 ± 0.0%	3.53 🕏 0.12	2.89 ± 0.1
PS-10 PS-20	54.0 ± 3.2	333.6 4 12.1	1(x).6 x 6.1	formers as rescribed	. No student and Mes	hade and in the log	end to Mg. 2

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flucturescence of a reporter die on the S' and. The degree of quenching is sufficient for this type of allgomicleotide to be used as a probe in the S' nuclease PCR acces.

To test the hypothesis that quanching by a 2' TAMBA depends on the flexibility of the oligonucloode, fluorescence was measured for probes in the singlestranded and double stranded stages. Tohis 3 reports the fluorescence observed at \$18 and \$82 nm. The relative degree of quenching is assessed by calculating the RQ ratio. For probes with TAMRA K-10 nucleotides from the 5' end, there is little difference in the RQ values when comparing single-stranded with doublestranded obsonucleotides. The results for probes with TAMPA at the 3' and are much different For these probes, bybridication to a complementary straind causes a dramatic increase in ItQ. We propose that this loss of quenching is caused by the rigid structure of double. stranded DNA, which prevents the 5 and 3' ends from being in proximity.

When TAMRA is placed toward the 3 and, there is a marked Mg<sup>2+</sup> effect on quenching. Figure 3 shows a plot of observed RQ values for the A1 series of probes as a function of Mg<sup>2+</sup> concentration. With TAMRA attached near the 5 and (probe A1-2 or A1-7), the RQ value at 0 mm Mg<sup>2+</sup> is only slightly higher than RQ at 10 mm Mg<sup>2+</sup>. For probes A1-19, A1-22, and A1-26, the RQ values at 0 mm Mg<sup>2+</sup> are very high, indicating 4 much

reduced quanching efficiency. For each of these probes, their is a marked docrease in MQ at I mm Mg2.1 fallowed by u gradual decline as the Mg2 cunceritrution increases to 10 mm. Probu A1-14 shows an intermediate RQ value at 0 mm Mg74 with a gradual decline at higher Mg24 concentiations, In a low-salt covironment with no Mga, present a singla-stranded oligonuclentide would be expected to adopt an extended conformation because of electrostatic repulsion. The binding of Mg2+ tons acts to shield the negative charge of the phosphate hackbone so that the oligoniucies otide can adopt conformations where the 3' end is close to the 5' end. Therefure, the observed Mg2+ effects support the notion that quenching of a 5' reporter dye by IAMRA at or near the 3' end depends on the flexibility of the oilgonucleodde.

### DISCUSSION

The striking finding of this study is that it seems the modamine dye TAMRA, placed at any position in an oligonucleotide, can quench the fluorescent emission of a fluorescent (6-FAM) placed at the Stender This implies that a single-stranded, double-laboted oligonucleotide must be able to adopt conformations where the TAMRA is close to the 6 end. It should be noted that the datay of 6-FAM in the excited state requires a certain amount of time. Therefore, what

matters for quenching is not the average distance between 6-FAM and TAMRA but, rither, how close TAMRA can get to 6-FAM during the lifetime of the 0-FAM excited state. As long as the ducay time of the excited state is relatively long compared with the molecular motions of the oligomucicotide, quenching can occur. Thus, we propose that TAMRA at the 3 end, or any other position, can quench 6-FAM at the 3' and because TAMRA is in proximity to 6-FAM often enough to be able to accept energy transfer from an excited 6-FAM.

Details of the fluorescence measurements remain puzeling, For example, Table 3 shows that hypridization of probes A1-26, A3-24, and 1/5-28 to their complementary strands not only course a large increase in 6-PAM fluorescence at \$18 nm but also causes a modest increase in T'AMRA fluorescence at 582 min. If TAMILA IS boing excited by energy transfor from quenched 6-FAM, then loss of quenching attributable to hybridization should cause a decrease in the fluorescence emission of TAMRA. The fact that the fluorescence emission of TAMRA Increases indicates that the situation is more complex. For example, we have anecumal evidence that the bases of the oligonuclectide, especially (i, quench the fluorescence of both 6-RAM and TAMPA to some degree. When doublestranded, base-pairing may reduce the ability of the bases to quench. The primany factor causing the quenching of 6-PAM in an intect probe is the TAMBA dyc. Pyldence for the Importance of TAMRA IS that O FAM Housescence remains relatively unchanged when probes lehaled only with 6-1/AM are used in the 5' nucleose PCR assay (data not shown). Secondary effectors of fluorescence, both before and after cleavage of the probe, need to be explored further.

Regardless of the physical mochanism, the relative independence of position and quenching greatly simplified the design of probes for the St muclease PCR assay. There are three main factors that determine the performance of a double-tabeled fluorescent probe in the St nuclease PCR assay. The first factor is the degree of quenching observed in the intest probe. This is characterized by the value of RQT, which is the ratio of reporter to quencher fluorescent emis-

TABLE 3 Comparison of Pharescence Endodous of Single-stranded and Double-stranded Puorogenic Profes

DODDIC-MINISTRACO LINGUISTA						
	518 nm		70g S82		RQ	
- احا	43	ds	93	1 de 2	16	d3
	37.76	P.A.A.	80.16	138,18	0,45	11.50
AT-7	2.7.75		53.50	93.86	0,84	5.43
A1.26	43.51	8E,908			0.43	0.38
ARIA	16.75	62.88	19,11	165.57		
A3-24	30.05	578.64	67,77	140128	(),45	3.23
		70.13	\$4.63	121.09	0,54	o.sg
C5-2	35.02			61.13	0,61	\$.25
1'2-27	20.R0	220.47	6\$.1U	· ·		
15-10	27,14	144.85	62.95	165.54	0.44	0.87
75-10 75-20	33.66	462.20	72.30	104.41	0.46	4.43

(45) Single-stranded, The fluorescence emissions at \$18 or \$82 nm for solutions containing a final concentration of \$0 nm indicated probe, 10 mm Tris-HCI (pH 8.5), 50 mm XCI, and 10 mm MgCl<sub>2</sub>, (ds) Double-stranded. The initiality contained, in addition, 100 nm A1C for probes A1-7 and A1-76, 100 nm A2C for probes A3-6 and A3-24, 100 nm F2C for probes F2-7 and F2-77, or 100 nm F3C for probes F3-10 and F3-28. Before the addition of MgCl<sub>2</sub>, 120 µL or early sample was neared

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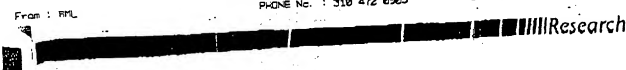
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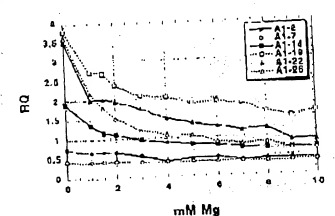


FIGURE 3 Lillest of Mg" concentration on RQ rado for the Al series of probes. The fluorostiente emission intendity at \$18 and \$82 nm was measured for solutions confeining 50 nm probe, 10 mm Tris-HCl (pH 8.3), 50 mm KCl, and varying amounts (0-10 mm) of MgCls. The calculated KCl ratios (818 mm lotensity divided by 342 nm limitally) are platfed vs. MgCls concentration (inm and the latensity divided by 342 nm limitally) are platfed vs. MgCls concentration (inm and limitally). Ma). The key (upper right) shows the probas examinal.

dyes used, specing between reporter and quencher dyes, nucleotide sequence context effects, presence of structure or uther factors that reduce fleatbility of the oligonucleotide, and purity of the probe. The second factor is the efficiency of hybridization, which depends on probe Time presence of secondary structure in probe or template, annealing emperature, and other exaction conditions. The third factor is the efficiency at which Tag DNA polymerase cleaves the bound probe between the reporter and quancher dyes. This cleavage is dependent on sequence complementarity between probe and template as shown by the observation that mismatches in the regulent between reporter and quencher dyes arestically reduce the thravake of hteps.(1)

The rise in RQ' values for the A1 senet of probes seems to indicate that the degree of quenching is reduced somewhat as the quencher is placed toward the 3' and. The lowest apparent quanthing is observed for probe A1-19 (san Fig. 3) rainer than for the probe where the TAMRA is at the 3' and (A1-26). This is understandable, as the conformation of the 3' end position would be expected to ha less restricted than the conformation of an internal position. In effect, a quencher at the &' and is freez to adopt conformations close to the 5' reporter dye than is an internally placed

probes, the interpretation of RCL values is less clearent. The All probes show the some (rend as A1, with the 3' TAMRA probe having a larger RQ" than the laternal TAMIN probe. For the P2 pali, pull probat have about the same RCI. value. For the PS probes, the RQ for the 3' probe is less than for the internally labeled probe. Another factor that may explain some of the observed variation is that purity affects the RQ value. Although all probes are HPLC putilied, a small amount of contamination with unquenched reporter can have a large effeet on NO

Although there may be a modest elfeel on degree of quenching, the posttion of the quencher apparently can linve a large effect on the efficiency of probe cleavage. The most drastic effect it observed with probe A1-2, where placement of the TAMRA on the second nocleative reduces the efficiency of cleanage to almost zoro. For the A3, I'2, and PS probes, ARQ is much greater for the 3' TAMKA probes as compared with the internal TAMRA probes. This is explained most casily by assuming that probes with TAMRA at the 3' and are more likely to be cleaved between reporter and quencher than are probes with TAMRA altached internally. Por the Al probes, the cleavage efficiency of probe A1-7 must already be quite high, as ARQ does not increase when the quencher is does to the 3' end. This illus-

trates the importance of hoing while to use probes with a quencher on the X' end in the 5' nucleuse I'CR unsay. In this assay, an increase in the intensity of reporter fluorescence is observed only when the probe is cleaved between the reporter and quencher dyes. By placing the repeater and adenotics dyes on the opposite ends of an oligonuclectide probe, any cleavage that occurs will be detected. When the quencher is attached to an incomest nucleotide, sometimes the probe works well (A1-7) and other times not so well (A3-6). The relatively pour performance of probe A3-6 presumably mesor the probe it being cleaved 3' to the quenchor rather than netween the reporter and quencher. Therefore, the best chance of having a probe that reliably detects accumulation of PCR product in the 5' nuclease FCR assay is to use a probe with the reporter and quencher tiyes on opposite ends.

Placing the quencher dye on the 3' and may also provide a stight bonefit in terms of hybridization efficiency. The presence of a quencher attached to an internal nucleatide might be expected to discapt base-pairing and reduce the Tm of a probe. In fact, a 201-101 reduction in T, has been observed for two probes with internally scanded TAMKAs. (\*\*) This disruptive effect would be minimized by placing the quencher at the 3' end. Thus, probes with I' quenchers might exhibit alightly higher hybridization efficiencies then probes with Internal quenchers.

The combination of increased cleavage and hybridisation efficiencies means mat probes with 3' quenebers probably will be more tolerant of mismatches between probe and target as compared with internally labeled probes. This tolerance of mismatches can be advantageous, as when trying to use a single probe to detect PCR-amplified products from samples of different species. Also, it means that cleavage of probe during PCR is less scribble to alterations in an. dealing remperature or other seaction conditions. The one application where tolerance of mismatches may be a disadvantage is for allelie discrimination. Lee et al (3) demonstrated that allele-specific probes were cleaved between reporter and quancher only when hybridized to z perfectly complementary larget. This allowed them to distinguish the normal human cystic fibrosis allele from the AF508 mutant. Their probes had TAMRA attached to the seventh nucleotlde from

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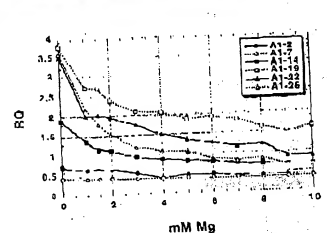


FIGURE 3. Effect of Mg<sup>61</sup> to resent ration on RQ ratio for the A1 series of probes. The fluorest states are summarised by 1518 and 182 nm was measured for solutions concluding 50 nm probe, 10 nm missing instances of MgCl<sub>2</sub>. The calculated RO Tris-IICI (pH 8.3), 50 nm KCl, and varying emounts (0.10 nm) of MgCl<sub>2</sub>. The calculated RO ratios (518 nm intensity divided by 582 nm intensity) are plotted vs. MgCl<sub>2</sub> concentration (mm Mg). The key (upper right) shows the probes examined.

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Placing the quenches dye on the 3' end may also provide a slight benefit in terms of hybridization efficiency. The presence of a quencher attached to an internal nucleotide anight be expected to discupt base-pairing and reduce the 7<sub>in</sub> of a probe. In fact, a 2°C-3°C reduction in T<sub>pin</sub> has been observed for two probes with internally attached TAMRAS. (9) This disruptive effect would be minimized by placing the quenchers at the 3' end. Thus, probes with 3' quenchers might exhibit alightly nigher hybridization efficiencies than probes with internal quanchers.

The combination of Increased cleavage and hybridization efficiencies means that makes with 3' quanchers probably will be more tolorant at mismaiches between probe and target as compared with internally labeled probes. This tolerance of mismatches can be advantagrous, as when trying to use a single probe to detect PCR-amplified products from samples of different species. Also, it means that cleavage of probe during PCK le less sensitive to alicrations in annealing tumperature or other reaction conditions. The one application where tolerance of mismarches may be a disadvantage is for allelic discrimination. Lec et al. (1) demonstrated that allele-specific bruties were cleaved petween saporter and quencher only when hybridized to a perfectly complementary ranges. This allowed them to distinguish the normal human cytic fibrosis allele from the AFSOR mutant. Their probes had TAMRA attached to the seventh nucleotide from From : BML

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# Researchilli

the 5' end and were designed to that any mismatches were between the reporter and quencher, increasing the distance between reporter and quencher would lessen the disruptive effect of mixmarches and allow cleavage of the probe on the incorrect target. Thus, probes with a quencher attached to an internal nucleotide may still be useful for allolic distrimination.

in this study lose of quenching upon hybridization was used to show that quenching by a 2' TAMRA is dependent on the flexibility of a single-erranded oilgonucleotide. The increase in reporter Ilvarascence Intensity, though, could also be used to determine whother bybridleation has accurred or not. Thus, oligonuclcouldes wills seporter and quenches dyes attached at opposite ends should also be useful as hybridization probes. The ability to detect hybridization in real time means that these probes could be used to measure hybridization kinetics. Also, this type of probe could be used to develop nomogeneous hybridiration essays for diagnostics or other applications. Bagwell et al. [10] describe just this type of hamogeneous assay where hybridization of a probe causes an incrussa in fluorosconco caused by a loss of quenching. However, they utilized a complex probe design that requires adding nucleatities to both ends of the probe requescy to form two imperfect hairpins. The tosults presented here demonstrate that the simple addition of a reporter dye to une end of an oligonius cleotide and a quencher dye to the other and generates a fluoregenic probe that con detect hybridisation or PCR amplification.

#### **ACKNOWLEDGMENTS**

We acknowledge Lincoln McRride of Perkin-Elmer for his support and encouragement on this project and Mitch Winnik of the University of Toronto for helpful discussions on time wolved fluorescence.

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Received December 30, 1994) accepted in revised form March 6, 1995.

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SIMULTANEOUS AMPLIFICATION AND DETECTION OF SPECIFIC DNA SEQUENCES

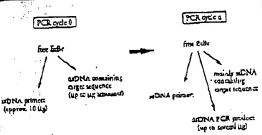
Russell Higuchi<sup>4</sup>, Gavin Dollinger<sup>1</sup>, P. Sean Walsh and Robert Grideric CA Finery III. CA P4008. 'Chiron Corporation, 1400 53rd St., Emeryville, CA P4008. 'Chiron Corporation, 1400 53rd St., Emeryville, CA P4008. 'Corresponding author. Russell Higuchi\*, Gavin Dollinger1, P. Sean Walsh and Robert Griffith

we have enhanced the polymerasa chain or the section (PCR) such that specific DNA reaction (PCR) such that specific DNA requences can be detected without opening the reaction tube. This enhancement requires the addition of ethidium bromide stranded (ds) DNA an increase in fluorescence in such a PCR indicates a positive range of externally. In fact, amplification can be continuously monitored in order to follow its progress. The ability to simultaneously amplifics and improves PCR and may facilitate its automation and more reduction and require straint between the simplification and may facilitate its automation and more straint progress and ethicity used in this string, even though it is four years since thermostable DNA polymeration of the processing steps, and blue positive results from the requirement of the string is done in order to determine whether the carget the finital base it to larget contributor to creat at the present thing is done in order to determine whether the carget the finite short in the string continuous in the contributor of creative divides of the processing once thermocytone form of "downstream" processing once thermocytone form of "downstream" processing once thermocytone on the processing of the processing in a risk of the proc

carryover" false positives in subsequent testing;

These downstream processing steps would be eliminated if specific amplification and desection of amplified DNA took place simultaneously within an unopened reaction vessel. Assays in which such different processes take place without the need to separate reaction components have been bernied homogeneous. No truly homogeneous PCR assay has been demonstrated to date, although progress towards this end has been reported. Chehab, et al. 1. developed a PCR product detection acheme using fluorescent primers that resulted in a fluorescent PCR product. Allelospecific primers, each with different fluorescent tags, were used to indicate the genotype of the DNA. However, the unincorporated primers must still be removed in a downstream process in order to visualize the result. Recontly, Holland, et al. 4, developed an assay in which the endogenous 5' exonuclease assay of Taq LINA polymerase was exploited to cleave a labeled oligonucleo-tide probe. The probe would only cleave if PCR amplitude carion had produced its complementary sequence. In order to detect the cleavage products, however, a subsequent process is again needed.
We have developed a truly homogeneous assay for PCR

and PCK product detection based upon the greatly increased Augrescence that ethidium bromide and other DNA binding dyes exhibit when they are bound to de-DNA 1-46. As outlined in Figure I, 2 prototypic PCR



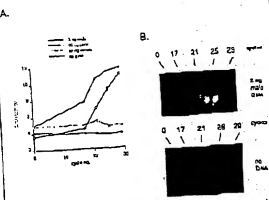
PCE 1 Principle of simulmoous amplification and detection of PCR product. The components of a PCR containing FLBY that are fluorescent are listed—EcBr itself, EtBr bound to either at DNA or dabna. There is a large fluorescence enhancement when Rift is bound to DNA and binding it gready caliance when DNA is double-stranded. After sufficient (n) cycles of PGR, the net increase in dabna results in additional Rift binding, and a net increase in total Augrescence

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PRIME 2 Cal electrophoresis of PCR amplification products of the human, nuclear gene, HIA DQa, made in the presence of increasing amounts of EdBr (up to 8 ug/ml). The presence of EdBr has no obvious effect on the yield or specificity of amplification.



RCUM 2 (A) Fluorescence measurements from PCRs that contain 0.6 µg/ml EtHr and that are specific for Y-chromosome report sequences. Five replicates PCBs were begun containing each of the DNA specified. At each indicated cycle, one of the five replicate PCRs for each DNA was removed from terrmocycling and its fluorescence measured. Units of fluorescence are arbitrary. (B) IV photography of PCR tubes (0.5 ml Eppendorf-style, polypropylene micro-contribuge tubes) containing reactions, those startpylene micro-contribute tubes (0.5 ml Eppendorf style, polypro-ing from 2 ng male 11NA and control reactions without any DNA, from (A).

begins with primers that are single-stranded DNA (sponsor), dNTPs, and DNA polymerase. An amount of diDNA containing the rarget sequence (target DNA) is also typically present. This amount can vary, depending on the application, from single-cell amounts of INA To on the application, from single-cell amounts of INA To on the application, if Ether is present, the reagents dias will Audrosce, in order of increasing Audrescence, are free EtBr isself, and RtBr bound to the single-stranded DNA primers and to the double-stranded target DNA (by its interculation between the stacked bases of the DNA double-helix). After the first denaguration cycle, target INA will be largely single-stranded. After a PCR is completed, the most significant change is the increase in the amount of daDNA (the YCR product icelf) of up to several micrograms. Formerly free EtBr is bound to the additional diDNA, resulting in an increase in fluores-cence. There is also some decrease in the amount of seDNA primer, but because the binding of ErBr to seDNA is much less than to dsDNA, the effect of this change on the total Huorescence of the sample is small. The fluoressonce increase can be measured by directing exchange illumination through the walls of the amplification vessel

before and after, or even continuously during, thetinocy.

#### RESULTS

PCR in the presence of Ethr. In order to assess the affect of EtBr in PCR, amplifications of the human HLA
DUn genero were performed with the dye present at
concentrations from 0.06 to 3.0 µg/ml (a typical concenconcentrations troid in saining of nucleic and following gel electrophoresis is 0.5 µg/m). As shown in Figure 9, gel electrophoresis revealed telle or no difference in the yield or quality of the amplification product whether Eth: was absent or present at any of these concentrations, indicating that ETAT does not inhibit PICR.

Detection of human Y-chromosome specific se-uences, Sequence-specific, fluorescence anhancement of quences. Sequence-spooning the demonstrated in a series of EtBr as a result of PCR was demonstrated in a series of amplifications containing 0.5 µg/ml FiBr and primers specific to repeat DNA sequences found on the human Y-chromosome of These PCRs initially contained either 60 ng male, 60 ng female, 2 ng male human or no DNA.
Five replicate Pt. Rs were begun for each DNA. After 0,
17, 21, 24 and 29 cycles of thermocycling, 2 PCR for each
17, 21, 24 and 29 cycles of thermocycling, 2 PCR for each DNA was removed from the thermocycler, and its fluorescence measured in a spectrofluorometer and plotted vs. amplification cycle number (Fig. 3A). The shape of this curve reflects the fact that by the time an increase in fluorescence can be detected, the increase in BNA is becoming linear and not exponential with cycle number. As shown, the Auorescence increased about three-told over the background Huorescence for the PCRs containing human male DNA, but did not significantly increase for negative control PCRs, which contained either no DNA or human female DNA. The mere male DNA present to begin with -60 ng versus 2 ng-the fewer cycles were needed to give a detectable increase in fluorescence. Gel electrophoresis on the products of these emplifications showed that DNA fragments of the exported size were made in the male DNA containing reactions and that little DNA synthesis took place in the control eamples.

In addition, the increase in Augrescence was visualized by simply laying the completed, unopened PCRs on a UV transilluminator and photographing them through a red filter. This is shown in figure 3E for the reactions that began with 2 ng male DNA and those with no DNA.

Detection of specific alleles of the human \$\textit{\theta}\circ{\theta}{\theta}\text{of the human } \text{\$\theta}\circ{\theta}{\theta}\text{of the human } \text{\$\theta}\text{of the human } \text{of the human }

Bene Iu order to demonstrate ther this abblioach pas adequate specificity to allow genetic ocreeming, a detection of the sickle-cell anemia mulation was performed. Figure 4 shows the fluorescence from completed amplifications containing EtBr (0.5 µg/ml) as detected by photography of the reaction tuber on a UV transilluminator. These reactions were performed using primers specific for er the wild-type or sickle-cell mutation of the human B-globin genera. The specificity for each allele is imparted by placing the sickle-mutation six at the terminal 3 nucleoide of one primer Byttsia as a presented and a property of the sickle-mutation six at the terminal 3 nucleoide of one primer Byttsia as a presented and a property of the six as a presented and a property of the six as a presented and a property of the six as a presented and a property of the six as a presented and a property of the six as a presented and a property of the six as a presented and a property of the six as a presented and a property of the six as a presented and a property of the six as a presented and a property of the six as a presented and a property of the six as a presented and a property of the six as a property of the nucleotide of one primer. By using an appropriate primer annealing temperature, primer extension—and thus any prince is complementary to the 8-globin allele present

Fach pair of amplifications shown in Figure 4 consists of reaction with either the wild-type allele special (left tube) or sickle-allele apscafic (right tube) primers. Parce different DNAs were typed: DNA from a homozygous wild-type \( \beta \)-globin individual (AA); from a heteroxygous sickle \( \beta \)-globin individual (AS); and from a homozygous sickle \( \beta \)-globin individual (AS). Each INA (50 c) general sickle \( \beta \)-globin individual (SS). Each INA (50 c) general sickle \( \beta \)-globin individual (SS). DNA to start each PCR) was analyzed in triplicate (3 pairs

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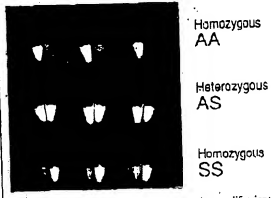
of reactions each). The DNA type was reflected in the relative fluorescence intensities in each pair of completed amplifications. There was a significant increase in fluoresence only where a  $\beta$ -globin allele DNA matched the printer set. When measured on a spectrofluorometer learning the shown the fluorescent and the set of the fluorescent and the fluorescent and the set of the fluorescent and the fluorescent and the set of the fluorescent and the set of the fluorescent and the set of the set of the fluorescent and the set of the (gats not shown), this fluoresoence was about three cimes (data not shown), this nuorescence was about three times that present in a PCR where both p-globin alleles were mismarched to the primer set. Cel electrophoresis (not shown) established that this increase in fluorescence was shown) due to the synthesis of nearly a microgram of a DNA fragment of the expected size for \$1-globin. There was little synthesis of deDNA in reactions in which the allelespecific primer was mismatched to both alletes:

Services of the services of th Condavous monitoring of a PCR Using a fiber optic device, it is possible to direct excitation illumination from a spectrofluorometer to a PCR undergoing thermocycling and to return is fluorescence to the spectrofluorometer. The Eurorescence readout of such an arrangement, directed at an EiBr-containing amplification of Y-chromosome specific sequences from 25 ug of human male DNA. is shown in Figure 5. The readout from a control PCR with no target DNA is also shown. Thirty cycles of PCR were menitored for each.

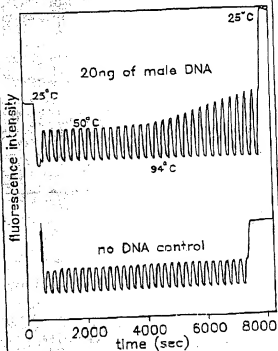
The fluorescence trace as a function of time dearly shows the effect of the thermocycling. Fluorescence intennicy rises and falls inversely with temperature. The fluoresecuce intensity is minimum at the denaturation comperature (94°C) and maximum at the annealing/extension temperature (50°C). In the negative-control POR, these fluorescence maxima and minima do not change signifieartly over the thirry thermocycles, indicading that there is Midde dsDNA synthesis without the appropriate target DNA, and there is little if any bleaching of FART during the continuous illumination of the sample.

in the PCR containing male DNA, the fluorescence maxima at the annealing extension temperature begin to increase at about 4000 seconds of thermocyching, and condute to increase with time, indicating that danNA is conduce to increase with ame, indicating that dsDNA is conduced at a detectable level. Note that the fluoring produced at a detectable level. Note that the fluoring increase minima at the denaturation temperature do not disminimately increase. Presumably because at this temperature of the amplification is followed by tracking the fluorescence increase at the annealing temperature. Analysis of the products of these two amplifications by gel electrophotes in the products of these two amplifications by gel electrophotes in the products of these two amplifications by gel electrophotes in the products of these two amplifications by gel electrophotes in the products of these two amplifications by gel electrophotes in male DNA containing sample and no detectable DNA indications. The products for the control sample.

Discussion by FOR The elimination of these processes the particle of the case of sickle-cell immalified decends solely on that of PCR is the case of sickle-cell immalified in the particle production of daDNA in the case of sickle-cell immalified in the production of daDNA in the absence of the production of daDNA in the absence of the production of other DNA that must be taken with the specificity required to descent pathogens can be detending on the number of pathogens in the sample and ic left. The specificity required to detect pathogens can be detending on the number of pathogens in the sample and ic left. The specificity required to detect pathogens can be detending on the number of pathogens in the sample and ic left. The specificity required to detect pathogens can be detending on the number of pathogens in the sample and ic left. The specificity required to detect pathogens can be supplied a difficult target is HIV, which requires detection tygous. The specificity is performed an electrophic containing at least tygous. The specificity is performed an electrophic containing at least tygous. The specificity is the specificity and the input of more total tygous. being produced as a desectable level. Note that the fluo-



REURI 4 UV photography of PCR tuber containing amplifications using EtBr that are specific to wild-type (A) or nickle (S) allelet of the human β-globin gene. The left of each pair of tuber contains allele-pecific primers to the wild-type allelets, the right tube primers to the nickle allele. The photograph was taken after 30 system of PCK, and the input DNAs and the alleles they contain are indicated. Fifty ug of DNA was used to begin PCR. Typing was done in triplicate (3 pairs of PCKs) for each input DNA.



PIEURI 5 Conditions, real time monitoring of a PCR. A fiber opic war used to carry excitation light to a PCR in progress and also emitted light back to a fluorouscher (see Experimental Protocol). Amplification using human male DNA specific primers in a PCR starting with 20 ng of human male DNA (top), or in a control starting with 20 ng of human male DNA (top), or in a control PCR without DNA (hontom), were monitored. There cycles of PCR were followed for each. The temperature cycled between 94°C (deuacuration) and 50°C (anacating and extension). Note in the mele DNA PCR, the cycle (time) dependent increase in fluorescence at the annealing/execution temperature.

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(\*) \*\*\* 1 / 37 **计图5单列**第 DNA-up to microgram amounts-in order to have suf-ficient numbers of ourget sequences. This large amount of starting DNA in an amplification significantly increases the background fluorescence over which any additional duoressence produced by PCR must be detected. An additional complication that occurs with targets in low copy-number is the formation of the "premar-dimer" artifact. This is the result of the extrusion of one primer using the other primer as a templare. Although this occurs infrequently, once it occurs the extension product is a substrace for PCR amplification, and can compose with true PCR targets if those targets are rare. The primer dimer product is of course dsDNA and thus is a potential

cource of false signal in this homogeneous assay.
To increase PCR specificity and reduce the effect of primer-dimer amplification, we are investigating a number of approaches, including the use of nested-primer amplifications that take place in a single tube, and the "hot-start", in which nonspecific amplification is reduced by raising the temperature of the reaction before DNA synthesis begins<sup>26</sup>. Preliminary results using these approaches suggest that primer-dimer is effectively reduced and it is possible to detect the incresse in first duorescence in a PCR instigated by a single HIV genome in a background of 10° colls. With larger numbers of cells, the background fluorescence contributed by genomic LINA becomes problematic. To reduce this background, it may be possible to use sequence-specific DNA-binding dyes that can be made to proferentially hind PCR product over genomic DNA by incorporating the dye-binding DNA sequence into the PCR product through a 5" "add.on" to the oligonucleoade primer

We have shown that the detection of fluorescence generated by an ErBr-containing PCR is straightforward, toth once PCR is completed and continuously during thermocyding. The aase with which automation of spacific IINA detection can be accomplished is the cross promising aspect of this assay. The fluorescence analysis of completed PCRs is already possible with existing instrumentation in 96-well format. In this format, the fluorescence in each PCR can be quantitated before, after, and even at selected points during thermocycling by moving the rack of PCRs to a 96-microwell plate fluorescence reader.

The instrumentation accessary to continuously monitor multiple PCRs simultaneously is also simple in principle. A direct octantion of the apparatus used here is to have multiple fiberoptics transmit the excitation light and flu-orescent emissions to and from multiple PCKs. The ability to menitor multiple PCRs continuously may allow quartindon of target DNA copy number. Figure 3 shows that the larger the amount of starting target DNA, the sooner during PCR a nuorescence increase is detected. Preliminary experiments (Higuchi and Dollinger, manuscript in preparation) with continuous monitoring have shown a sensitivity to two-fold differences in initial carget DNA concentration.

Conversely, if the number of target molecules is known—as it can be in genetic acreening—continuous monitoring may provide a meant of detecting false positive and false negative results. With a known number of Erzet molecules, a true positive would exhibit detectable Sucressence by a predictable number of cycles of PCK. Increases in fluorescence detected before or after that cycle would indicate potential artifacts. False negative results due to, for example, inhibition of DNA polymerase, may be detected by including within each PCR an inefficiently amplifying marker. This marker results in a fluorescence increase only after a large number of cycles-many more than are necessary to detect a true

positive. If a sample fails to have a fluorescence increase after this many cycles, inhibition may be suspected. Since, after this many cycles, inhibition may be suspected. Since, in this assay, conclusions are drawn based on the presence or absence of fluorescence cignal alone, such controls may be important. In any event, before any test based on this principle is ready for the clinic, an assessment of its false positive/false negative rates will need to be obtained uring a large number of known samples.

In summary, the inclusion in PCR of dyes whose huoreternce is enhanced upon binding daDNA makes it possible to detect specific DNA amplification from outside the PCR tube. In the future, instruments based upon this principle may facilitate the more widespread use of PCR in applications that demand the high throughput of

EXPERIMENTAL PROTOCOL

Muman HLA-DQa rene amplifications containing Ethn
PCK1 were set up in 100 µl volumes containing 10 mM Tra-PCG.

pH 8.3: 50 mM ECl; A mM McCl; 2.6 units of Iaq DNA
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ph 8.3: 50 mM ECl; A mM McCl; 2.6 units of Iaq DNA
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volume) cuntaining 0.5 pc/ml 2.63r werte prepared as described.
2. Thermocycling value of the proceeding primars and target (DNAs)
2. Thermocycling was 94°C for I min and 80°C for 1 min and 80°C for 1 min using 2 step-cycle program. The number of cycles for
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a cample were as indicated in Figure 3. Fluorescence measurement is described below.

Allele-specific, human a globin gene PCR. Amplifications of all closepocific, human a globin gene PCR. Amplifications of the property of

that was heteroxygous for the sickle trait (S), DNA that was heteroxygous for the sickle trait (AS), or DNA that was homezygous for the w.t. globin (AA). Thermusyting was for 30 keyetes at 94°C for 1 min. using a "seep-cycle" growth at 12° to provide allele-specific amplification. Completed Will et al. 2° to provide allele-specific amplification. Completed Will et al. 2° to provide allele-specific amplification. Completed Will et al. 2° to provide allele-specific amplification. Completed Will et al. 2° to provide allele-specific amplification. Completed Will et al. 2° to provide a red filter (Wrauen 1844). PCFs were photographed through a red filter (Wrauen 1844). PCFs were photographed through a red filter (Wrauen 1844). In some (UV-producus San Gabriel, CA).

Fluorespecific measurement. Finorespecific masulement were made on PCRs. Londinging Riffs in a Yluoruber 2° Buaranteet. (SPEX, Edison, N). Excitation was at the 500 nm band with (SPEX, Edison, N). Excitation was at the 500 nm band with (SPEX, Edison, N). Excitation was at the 500 nm band with (SPEX, Edison, N). Excitation was at the 500 nm band with (SPEX, Edison, N). Excitation was at the 500 nm band with (SPEX, Edison, N). Excitation was at the following the continuous allocation of the fiber was used used used to remove the excitation filter (Mallies allows). Continuous distributions of VCR. Continuous Continuous Continuous Continuous distributions of VCR. Continuous C

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vas used and the emission signal was radoed to the excitation signal to control for changes in light-source internal? Data were collected using the doubtood, version 2.6 (SPEX) data system.

"Years A Jed Streeting we thank Bob Jones for help with the spectroffuosmotric was thank Bob Jones for help with the spectroffuosmotric measurements and Heatherbell Tong for educing this manuscript.

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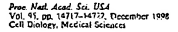
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## WISP genes are members of the connective tissue growth factor family that are up-regulated in Wnt-1-transformed cells and aberrantly expressed in human colon tumors

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Contributed by David Botstein and Arnold J. Levine, October 21, 1998

Wat family members are critical to many developmental processes, and components of the Wat signaling pathway have been linked to tumorigenesis in familial and sporadic colon carcinomas. Here we report the identification of two zeacs, WISP-1 and WISP-2, that are up-regulated in the mouse mammary epithelial cell line C57MG transformed by Wnt-1, but not by Wnt-4. Together with a third related gene, WISP-3, these proteins define a subfamily of the connective Ussue growth factor family. Two distinct systems demon-Strated WISP Induction to be associated with the expression of Wat-1. These included (i) CS7MG cells infected with a Wat-1 retrovirst vector or expressing Wnt-1 under the control of a tetracyline repressible promotor, and (ii) Wnt-1 transgenic mice. The WISP-I gene was localized to human chromosome 8q24.1-8q24.3. WISP-I genomic DNA was amplified in colon cancer cell lines and in human colon tumors and its RNA overexpressed (2- to >30-fold) in 84% of the tumors examined compared with patient-matched normal mucosa. WISP-J mapped to chromosome 6q12-6q23 and also was overex-pressed (4- to > 40-fold) in 63% of the colon tumors analyzed. In conteast, WISP-2 mapped to human chromosome 20u12-20q13 and its DNA was amplified, but RNA expression was reduced (2- to >30-fold) in 79% of the tumors. These results suggest that the WISP genes may be downstream of Wnt-1 signaling and that aberrant levels of WISF expression in colon cancer may play a role in colon tumorigenesis.

Wat-1 is a member of an expanding family of cysteine-rich, glycosylated signaling proteins that mediate diverse developmental processes such as the central of cell proliferation, adhesion, cell polarity, and the establishment of cell fates (1, 2). Wnt-1 originally was identified as an oncogone activated by the insertion of mouse mammary tumor virus in virus-induced mammary adenocarcinomas (3, 4). Although Wnt-1 is not expressed in the normal mammary gland, expression of Wnt-1 in transgenic mice causes mammary tumors (5).

In mammalian cells. Wat tamily members initiate signaling by binding to the seven-transmembrane spanning Frizzled receptors and recruiting the cytoplasmic protein Dishevelled (Dsh) to the cell memorane (1, 2, 6). Dsh then inhibits the kinase activity of the normally constitutively active glycogen synthase kinace-3\$ (GSK-3\$) resulting in an increase in B-catenin levels. Stabilized B-catenin interacts with the transcription factor TCF/Lefl, forming a complex that appears in the nucleus and binds TCF/Left target DNA elements to activate transcription (7, 8). Other experiments suggest that the adenomatous polyposis coli (APC) tumor suppressor gene also plays an important role in Wat signaling by regulating  $\beta$ -catenin levels (9). APC is phosphorylated by GSK-3 $\beta$ , binds to B-catonin, and facilitates its degradation. Mutations in either APC or \(\beta\)-catonin have been associated with colon carcinomas and melanomas, suggesting these mutations contribute to the development of these types of unneer, implicating

the Wat pathway in tumorigenesis (1).

Although much has been learned about the Wat signaling pathway over the past several years, only a few of the transcriptionally activated downstream components activated by Wat have been characterized. Those that have been described cannot account for all of the diverse functions attributed to Wat signaling. Among the candidate Wat turget genes are those encoding the nodal-related 3 gene, Kard, a member of the transforming growth tactor (TGF)- is superfamily, and the homeobox genes, engrailed, goosecoid, min (Xtwn), and siamois (2). A recent report also identifies r-myc as a target gene of the

(2). A recent report asso trentities to myc as a target gene of the Whit signaling pathway (10).

To Identify additional downstream genes in the Whit signaling pathway that are relevant to the transformed cell phenotype, we used a PCR-based cDNA subtraction strategy, suppression subtractive hybridization (SSH) (11), using KNA isolated from C17MO mouse manimary epithelial cells and C17MO cells stably transformed by a Whit-I certovirus. Overgression of Whit-I in this cell line is sufficient to induce a expression of Wat-I in this cell line is sufficient an induce a partially transformed phenotype, characterized by elongated and refractile cells that lose contact inhibition and form a multileyered array (12, 13). We reasoned that genes differentially expressed between these two cell lines might contribute to the transformed phenotype.

In this paper, we describe the cloning and characterization of two genes up-regulated in Wat-1 transformed cells, WISP-1 and WISP-2, and a third related gene, WISP-3. The WISP reno are members of the CCN family of growth factors, which includes connective tissue growth factor (CTGF), Cyr61, and nov, a family not previously linked to Whit signaling.

### MATERIALS AND METHODS

SSH. SSH was performed by using the PCR-Salect cDNA Subtraction Kit (CLONTECH). Tester double-stranded

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Anhreviations: TCF, transforming growth factor: CTCF, connective Association for transforming growth factor. The connective five time to the connective five time to the connective five time the paper factor type C module.

Data deposition: The sequences reported in this paper have been deposited in the Genbank database (accession nos. AF100777, AF100779, AF100780, and AF100781).

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Proc. Natl. Acad. Sci. USA 95 (1958)

cDNA was synthesized from 2 µg of poly(A)<sup>+</sup> RNA isolated from the C5/MG/Wnt-1 cetl line and driver cDNA from 2 µg of poly(A)<sup>+</sup> RNA from the parent C57MG cells. The subtracted cDNA library was subcloned into a pGEM-T vector for further analysis.

cDNA Library Screening. Clones encoding full-length mouse WISP-1 were isolated by screening a Agt10 mouse embryo cDNA library (CLONTECH) with a 711-bp probe from the original partial clone 568 saquence corresponding to amino acids 128-169. Clones encoding full-length human WISP-1 were isolated by screening Agt10 lung and fetal kidney cDNA libraries with the same probe at low stringency. Clones encoding full-length mouse and human WISP-2 were isolated by screening a C57MG/Wnt-1 or human fetal lung cDNA library with a probe corresponding to nucleocides 1463-1512. Full-length c17NAs encoding WISP-3 were cloned from human bone marrow and fetal kidney libraries.

Expression of Human WISP RNA. PCR amplification of first-strand cDNA was performed with human Multiple Tiesue cDNA panels (CLON1ECH) and 300 µM of each dNTP at 94°C for 1 sec, 62°C for 30 sec, 72°C for 1 min, for 22-32 cycles. WISP and glyceraldshyde-3-phosphate dehydrogenase primer sequences are available on request.

In Site Hybridization. DP-labeled sense and antisense riboprobes were transcribed from an \$97-bp PCR product corresponding to nucleotides 601-1440 of mouse WISP-1 of a 294-bp PCR product corresponding to nucleotides 62-373 of mouse WISP-2. All tissues were processed as described (40).

Radiation Hybrid Mapping. Genomic DNA from each hybrid in the Stanford G3 and Genebridge 4 Radiation Hybrid Panels (Research Genetics, Huntsville, AL) and human and hamster control DNAs were FCR-amplified, and the results were submitted to the Stanford or Massachusetta Institute of Tachnology web servers.

Cell Lines, Tumors, and Mucosa Specimens. Tissue specimens were obtained from the Department of Pathology (University of Pittsburgh) for patients undergoing colon resection and from the University of Locds. United Kingdom. Genomic DNA was isolated (Qiagen) from the profied blood of 10 normal human donors, surgical specimens, and the following ATCC human cell lines: SW480, COLO 320DM. HT-29, WiDr, and SW403 (colon adenocarcinomas). SW620 (lymph node motaetasis, colon adenocarcinoma), HCT 116 (colon carcinoma), SK-CO-1 (colon adenocarcinoma, ascites). 2nd HM7 (a variant of ATCC colon adenocarcinoma cell line US 174T). DNA concentration was determined by using Hoochst oye 33258 intercalation fluorimetry. Total RNA was prepared by homogenization in 7 M GuSCN followed by centrifugation over CSCI cushlons or prepared by using RNAzol.

Gene Amplification and RNA Expression Analysis. Relative gane amplification and RNA expression of WISPs and c-mye in the cell lines, colorectal tumore, and normal mucosa were determined by quantitative PCR. Geno-specific primers and fluorogenic probes (sequences available on request) were designed and used to amplify and quantitate the genes. The relative gene copy number was derived by using the formula 2iad) where ACt represents the difference in amplification cycles required to detect the WISP genes in peripheral blood lymphosyte DNA compared with colon tumor DNA or colon tumor RNA compared with normal mucosal RNA. The o-method was used for calculation of the SE of the gene copy number or RNA expression level. The WISF-specific signal was normalized to that of the glyceraldehyde-3-phosphate dehydrogenase housekeeping gone. All TaqMan assay reagents were obtained from Perkin-Elmer Applied Biosystems.

#### RESULTS

Isolation of WISP-1 and WISP-2 by SSH. To identify Wnt-1-Inducible genes, we used the technique of SSH using the

mouse mammary epithetial cell line C57MO and C57MG cells that stably express Wnt-1 (11). Candidate differentially expressed eDNAs (1,384 total) were sequenced. Thirty-nine percent of the sequences matched known genes or homologues, 32% matched expressed sequence tags, and 29% had no match. To confirm that the transcript was differentially expressed, semiquantitative reverse transcription-PCR and Northern analysis were performed by using mRNA from the C57MG and C57MG/Wnt-1 cells.

Two of the oDNAs, WISP-1 and WISP-2, were differentially expressed, being induced in the C57MG/Wnt-1 cell line, but not in the parent C57MG cells or C57MG cells overexpressing Wnt-4 (Fig. 1.A and B). Wnt-4, unlike Wnt-1, does not induce the morphological transformation of C57MG cells and has no effect on \(\mu\$-catenin levels (13, 14). Expression of WISP-1 was up-regulated approximately 3-fold in the C57MG/Wnt-1 cell line and WISP-2 by approximately 3-fold by both Northern analysis and reverse transcription-PCR.

An indopendent, but similar, system was used to examine WISP expression after Wnt-1 induction. CSTMG cells expressing the Wnt-1 gene under the central of a tetracyclinerepressible promoter produce low amounts of Wnt-1 in the repressed state but show a strong induction of War-1 mRNA and protein within 24 hr after retracycline removal (8). The levels of Wat-1 and WISP RNA isolated from these cells at various times after tetracycline reminal were assessed by quantitative PCR. Strong induction of Wnt-1 mKNA was seen as early as 10 hr after tetracycline removal. Induction of WISP mRNA (2- to 6-fold) was seen at 48 and 72 hr (data not shown). These data support our previous observations that show that WISP induction is correlated with Wat-1 expression. Because the induction is slow, occurring after approximately 48 hr, the induction of WISPs may be an indirect response to Wat-1 signaling.

cDNA clones of human WISP-1 were isolated and the sequence compared with mouse WISP-1. The cDNA sequences of mouse and human WISP-1 were 1,766 and 2,830 bp in length, respectively, and encode proteins of 367 an, with predicted relative molecular masses of ~40,000 (M<sub>r</sub> 40 K). Both have hydrophobic N-terminal signal sequences, 38 conserved everine residues, and four potential N-linked glycosylation sites and are 84% identical (Fig. 24).

Pull-length cDNA clones of mouse and human WISP-2 were 1.734 and 1.293 bp in length, respectively, and encode proteins of 251 and 2.50 as, respectively, with predicted relative molecular masses of ~77.000 (M, 27 K) (Fig. 2H). Mouse and human WISP-2 are 13% identical. Human WISP-2 has no potential N-linked glycosylation sites, and mouse WISP-2 has one at

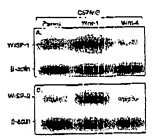


Fig. 1. WISP-I and RISP-2 are induced by Whit-I, but not Whit-I, expression in CIMC cells. Northern analytic of WISP-I (A) and WISP-2 (B) expression in CIMC, CIMC, CIMC/Whit-I, and CIMC/Whit-I cells. Poly(A)\* RNA (2 µg) was subjected to Northern blot analysis and hybridized with a 70-bp mouse WISP-I-specific probe (amino acids 173-300) or a 100-bp WISP-I-specific probe (aucleotides 1438-1627) in the Y untranslated teglon. Blots were rehybridized with human H-ectin probe.

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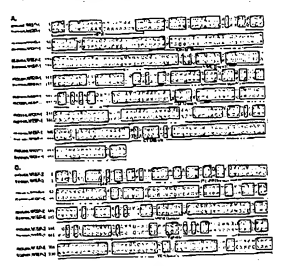


Fig. 2. Encoded amino acid sequence alignment of mouse and human WISP-1 (A) and mouse and human WISP-2 (B). The potential eignal sequence insulin-like growth factor-binding protein (15P-BP), www. thrombospondin (TSP), and C-terminal (CT) domains are underlined.

position 197. WISP-2 has 28 cysteine residues that are conserved among the 38 cysteines found in WISP-1.

Identification of WISP-3. To search for related proteins, we screened expressed asquence tag (EST) natabases with the WISP-1 protein sequence and identified several ESTS as potentially related sequences. We identified a homologous protein that we have called WISP-3. A full-length human WISP-3 cDNA of 1,371 hp was isolated corresponding to those ESTS that encode a 334-na protein with a prodicted molecular mass of 39,293. WISP-3 has two potential N-linked glycosylation sites and 36 cysteine residues. An alignment of the three human WISP proteins shows that WISP-1 and WISP-3 are the most similar (42% identity), whereas WISP-2 has 37% identity with WISP-1 and 32% identity with WISP-3 (Fig. 34).

WISPr Are Homologous to the CTCF Family of Proteins. Human WISP-1, WISP-2, and WISP-3 are novel sequences: however, mouse WISP-1 is the same as the recently identified Elm.) gene. Elm.1 is expressed in low, but not high, metastatic mouse melanoma cells, and suppresses the in vivo growth and metestatic potential of K-1735 mouse melanome colls (15). Human and mouse WISP-2 are homologous to the recently described rat gone, rCop-1 (16). Signuticant homology (36-44%) was seen to the CCN family of growth lactors. This family includes three members, CTGF, Cyr61, and the protooncogene nov. CTGF is a chemotectic and mitogenic factor for fibroblasts that is implicated in wound healing and abrotic disorders and is induced by TOF-8 (17). Cyr61 is an extracelfular matrix eignating molecule that premotes cell adhesion. proliferation, migration, angiogenesis, and tumor growth (18, 19). nov (nophroblastoma overexpressed) is an immediate early gene associated with quiescence and found altered in Wilins tumors (20). The proteins of the CCN family share functional, but not sequence, similarity to Wnr-1. All are secreted, cysteine-rich hoparin binding glycoproteins that associate with the cell surface and extracellular matrix.

WISP proteins exhibit the modular architecture of the CCN family, characterized by four conserved cysteine-rich domains (Fig. 3B) (21). The N-terminal domain, which includes the first 2 cysteine residues, contains a consensus sequence (GCGC-CXXC) conserved in most insulin-like growth factor (IGF)-

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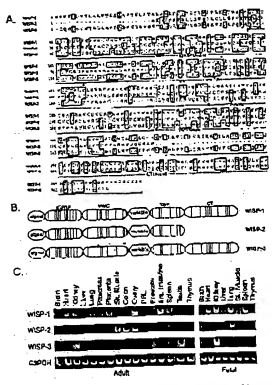


Fig. 3. (A) Encoded amino acid sequences alignment of human WISPs. The cysteine residuse of WISP-1 and WISP-2 that are not present in WISP-3 are indicated with a dot. (A) Schematic representation of the WISP proteins showing the normalin structure and cyrotine residuaes (varical lines). The four cysteine residuaes in the VWC domain that are absent in WISP-3 are indicated with a dot. (C) Expression of WISP mRNA in human tissues. PCR was performed on human multiple-assue cDNA panels (CLONTECH) from the indicated adult and fertal tissues.

binding proteins (BP). This sequence is conserved in WISP-2 and WISP-3, whereas WISP-1 has a glutamine in the third position instead of a glycine. CTCF recently has been shown to specifically bind IGF (22) and a truncated nov protein tacking the IGF-BP domain is oncogenic (23). The von Willebrand factor type C module (VWC), also found in certain collagens and mucins, covers the next 10 cysteine residues, and is thought to participate in protein complex formation and oligomerization (24). The VWC domain of WISP-3 differs from all CCN family memoers described previously, in that it contains only six of the 10 cysteine residues (Fig. 3 A and #). A short variable region follows the VWC domain. The third module, the thrombosopoudin (TSP) domain is involved in binding to sulfated glycoconjugates and contains the cysteins residues and a conserved WSxCSxxCG moul first identified in thrombospondin (25). The C-terminal (CT) module containing the remaining 10 cysteines is thought to be involved in dimerization and receptor binding (26). The CT domain is present in all CCN tamity members described to date but is absent in WISP-2 (Fig. 3 A and B). The existence of a putative signal sequence and the absence of a transmembrane domain suggest that WISPs are secreted proteins, an observation supported by an analysis of their expression and secretion from mammalian cell and baculovirus cultures (data not shown).

Expression of WISP mRNA in Human Tiesues. Tissuespecific expression of human WISPs was characterized by PCK 14720 Cell Biology, Medical Sciences: Penaica et al.

analysis on adult and fetal multiple tissue cDNA panels. WISP-1 expression was seen in the adult heart, kidney, lung, panereas, placenta evary, small intestine, and spleen (Fig. 3C). Little or no expression was detected in the brain, liver, skeletal muscle, colon, peripheral blood leukocytes, prostate, testis, or thymus. WISP-2 had a more restricted tissue expression and was detected in adult skeletal muscle, colon, evary, and fetal lung. Predominant expression of WISP-3 was soon in adult kiditey and testis and fetal kidney. Lower levels of WISP-3 expression were detected in placenta, ovary, prostate, and small intestine.

In Slau Localization of WISP-1 and WISP-2. Expression of WISP-1 and WISP-2 was assessed by in size hybridization in mammary rumors from Wnt-1 transgenic mice. Strong expression of WISP-1 was observed in stromal fibroblasts bring within the fibrovascular rumor stroma (Fig. 4 A-D). However, low-level WISP-1 expression also was observed focally within tumor cells (data not shown). No expression was observed in normal breast. Like WISP-1, WISP-2 expression also was seen in the tumor stroma in breast rumors from Wnt-1 transgenic animals (Fig. 4 E-H). However, WISP-2 expression in the stroma was in spindle-shaped cells adjacent to capillary vessels, whereas

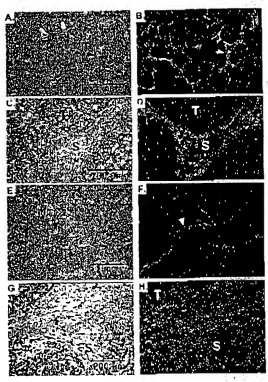


Fig. 4. (A.C.E. and O) Representative hematoxylin/cosin-stained images from breast tumors in Wnt-1 transpeals mice. The corresponding deck-field images showing WISP-1 expression are shown in B and D. The tumor is a moderately well-differentiated adenoceroinoma showing avidence of adenoid cytic change. At low power (A and B), expression of WISP-1 is seen in the delicate branching fibroviscular tumor stroma (arrowhead). At higher magnification, expression is seen in the stromal(s) fibroblasts (C auld D), and tumor cells are negative. Focal expression of WISP-1, however, was observed in tumor cells in some areas. Images of WISP-2 expression are shown in E-H. At low power (S and F), expression of WISP-1 is seen in cells lying within the fibroviscular tumor stroma. At higher magnification, those cells are negative (G and H).

the predominant cell type expressing WISP-1 was the stromal fibroblasts.

Chromosome Localization of the WISP Genes. The chromosomal location of the human WISP genes was determined by radiation hybrid mapping panels. WISP-1 is approximately 3.48 cR from the meiotic marker AFM259xc5 [logarithm of ours (lod) score 16.31] on chromosome 8q24.1 to 8q24.9, in the same region as the human locus of the novH femily member (27) and roughly 4 Mbs distal to e-myc (28). Preliminary fine mapping indicates that WISP-1 is located near D8S1712 STS. WISP-2 is linked to the marker SHGC-3.3922 (lod = 1,000) on chromosome 20q12-20q13.1. Human WISP-3 mapped to chromosome 6q22-6q23 and is linked to the marker AFM2112eS (lod = 1,000). WISP-3 is approximately 13 Mbs proximal to CTGF and 23 Mbs proximal to the human collular oncogene MYB (27, 29).

Amplification and Aberrant Expression of WISPs in Human Colon Tumors. Amplification of protococcogenes is seen in many human tumors and has citological and prognostic significance. For example, in a variety of tumor types, e-myc amplification has been associated with intalignent progression and poor prognosis (30). Because WTSP-1 resides in the same general chromosomal location (8q24) as c-myc. we asked whether it was a target of gene amplification, and, if so, whother this amplification was independent of the conyc locus. Genomic DNA from human colon cancer cell lines was assessed by quantitative PCR and Southern blot analysis. (Fig. 5 A and B). Both methods detected similar degrees of WISP-1 amplification. Most cell lines showed significant (2- to 4-fold) amplification, with the HT-29 and Willr cell lines demonstrating an 8-fold increase. Significantly, the pattern of amplification observed did not correlate with that observed for c-mye, indicating that the e-mye gene is not part of the amplicon that involves the WISA I locus.

We next examined whether the WISP genes were amplified in a panel of 25 primary human colon adenocarcinomas. The relative WISP gene copy number in each colon curnor DNA was compared with pooled normal DNA from 10 donors by quantitative PCR (Fig. 6). The copy number of WISP-1 and WISP-2 was rignificantly greater than one, approximately 2-fold for WISP-1 in about 60% of the tumors end  $\frac{1}{2}$  to 4-rold for WISP-2 in 92% of the tumors (P < 0.001) for each). The copy number for WISP-3 was indistinguishable from one (P = 0.166). In addition, the copy number of WISP-2 was significantly higher than that of WISP-1 (P < 0.001).

The levels of WISP transcripts in RNA isolated from 19 adenocarcinomas and their matched normal mucosa were

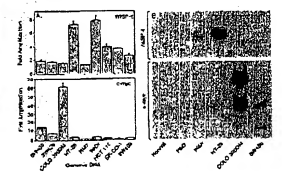


Fig. 5. Amplification of WISP-1 genomic DNA in colon cancer cell lines (A) Amplification in cell line DNA was determined by quantitative PCR. (B) Southern blots containing genomic DNA (10 µg) digister with FacR1 (WISP-1) or Abal (c-myc) were hybridized with a 100-bp human WISP-1 probe (amino acids 186-219) or a human compc probe (located at bp 1901-2000). That WISP and tage genes are dotted in normal human genomic DNA after a longer film exposure.

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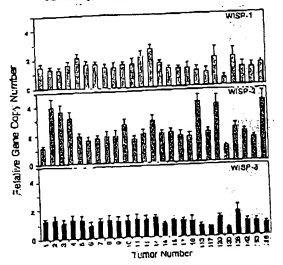


FIG. 6. Genomic amplification of WISP genes in human colon tumors. The relative scine copy number of the WISP genes in 25 adenocarcinomas was assayed by quantitative PCR, by cumpariae DNA from primary human tumors with pooled DNA from 10 healthy denors. The data are means ± SEM from one experiment done in triplicate. The experiment was repeated at least three times.

assessed by quantitative PCR (Fig. 7). The level of WISP-1 RNA present in tumor tissue varied but was significantly increased (1- to >25-fold) in 84% (16/19) of the human colon tumors examined compared with normal adjacent nucosa. Four of 19 tumors showed greater than 10-fold overexpression. In contrast, in 79% (15/19) of the tumors examined, WISP-2 KNA expression was significantly lower in the tumor than the mucosa. Similar to WISP-1, WISP-3 RNA was overexpressed in 63% (12/19) of the colon rumors compared with the normal

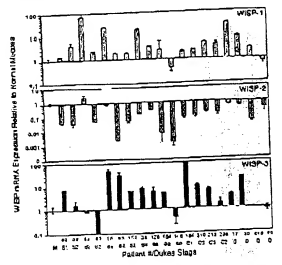


Fig. 7. WISP RNA expression in primary human color tumors relative to expression in normal mucosa from the same patient. Expression of WISP mRNA in 19 adenocarcinomas was assayed by quantitative FCR. The Dukes stege of the tumor is listed under the sample number. The data are means n SEM from one experiment done in triplicate. The experiment was repeated at least twice.

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mucosa. The amount of overexpression of WISP-3 ranged from 4- to >40-fold

### DISCUSSION

One approach to understanding the molecular basis of cancer is to identify differences in gene expression between concer cells and normal cells. Strategies based on assumptions that steady-state mRNA levels will differ between normal and mallgnant cells have been used to clone differentially expressed genes (31). We have used a PCR-based selection strategy. S3HL to identify genes selectively expressed in C57MG mouse mammary crithelial cells transformed by Wut-l.

Three of the genes isolated, WISP-1, WISP-2, and WISP-3, are members of the CCN family of growth factors, which includes CfGF, Cyth1, and nov, a family not previously linked to Wnt signaling.

Two independent experimental systems demonstrated that WISP induction was associated with the expression of Wnt-1. The first was C57MC cells infected with a Wnt-1 terroviral vector or C57MC cells expressing Wnt-1 under the control of a tetracylina-repressible promoter, and the second was in Wnt-1 transcenic mice, where breast tissue expresses Wnt-1, whereas normal breast tissue does not. No WISP RNA expression was detected in mammary tumors induced by pnlyoma virus middle T entigon (data not shown). These data suggest a link between Wnt-1 and WISPs in that in these two situations. WISP induction was correlated with Wnt-1 expression.

It is not clear whether the WISPs are directly or indirectly induced by the downstream components of the Wat-1 signaling pathway (i.e., \(\theta\)-catenin-TCF-1/Left). The increased levels of WISP RNA were measured in Wat-1-transformed cells, hours or days after Wat-1 transformation. Thus, WISP expression could result from Wat-1 signaling directly through \(\theta\)-catenin transcription factor regulation or alternatively through Wat-1 signaling turning on a transcription factor, which in turn regulates WISPs.

The WISPs define an additional subfamily of the CCN family of growth factors. One striking difference observed in the protein sequence of WISP-2 is the absence of a CT domain, which is present in CTGF, Cyr61, non, WISP-1 and WISP-3. This domain is thought to be involved in receptor binding and dimerization. Growth factors, such as 1 OF-1, platelet-derived growth factor, and nerve growth factor, which contain a cystine knot motificate as dimers (32). It is tempting to speculate that WISP-1 and WISP-3 may exist as dimers, whereas WISP-2 exists as a monomer. If the CT domain is also important for receptor binding. WISP-2 may bind its receptor through a different region of the molecule than the other CCN family members. No specific receptors have been identified for CTGF or now. A recent report has shown that integrin  $\alpha_i \beta_i$  serves as an adhesion receptor for Cyr61 (33).

The strong expression of WISP-1 and WISP-2 in cells lying within the fibrovascular tumor stroma in breast tumors from Wnt-1 transgenic animals is consistent with provious observations that transcripts for the related CTGF gene are primarily expressed in the tibrous stroma of manusary tumors (34). Epithelial cells are thought to control the proliferation of connective tissue stroma in manusary tumors by a cascade of growth factor signals similar to that controlling connective tissue formation during wound repair. It has been proposed that manusary tumor cells or inflammatory cells at the tumor interstitlal interface secrets TGF-β1, which is the stimulus for stromal proliferation (34). TGF-β1 is secreted by a large percentage of malignant breast tumors and may be one of the growth factors that stimulates the production of CTGF and

WISPs in the stroma.

It was of interest that WISP-1 and WISP-2 expression was observed in the stromal cells that surrounded the tumor cells

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(epithetial cells) in the Wnt-1 transgenic mouse sections of breast tissue. This finding suggests that paracrino signaling could occur in which the stromal cells could supply WISP-1 and WISP-2 to regulate tumor cell growth on the WISP extracelluler matrix. Stromal coll-derived factors in the extracellular matrix have been postulated to play a role in tumor cell migration and proliferation (35). The localization of WISP-1 and WISP-2 in the stromal cells of breast lumors supports this peracrine model.

An analysis of WISP-I gens amplification and expression in human colon tumors showed a correlation between DNA amplification and overexpression, whereas overexpression of WISP-3 RNA was seen in the absence of DNA amphitication. In contrast, WISP-2 DNA was amplified in the colon tumors, but its mRNA expression was significantly reduced in the majority of rumors compared with the expression in normal colonic mucosa from the same patient. The gene for human WISP-2 was localized to chromosome 20q12-20q13, at a region frequently amplified and associated with poor prognosis in node negative breast cancer and many colon cancers, suggesting the existence of one or more oncogenes at this locus (36-38). Because the center of the 20q13 amplicon has not yet boon identified, it is possible that the apparent amplification observed for WISP-2 may be caused by another general this

A recent manuscript on rCop-1, the ret orthologue of W15P-2, describes the loss of expression of this gene after cell transformation, suggesting it may be a negative regulator of growth in cell lines (16). Although the mechanism by which wrsp-2 RNA expression is down-regulated during malignant transformation is unknown, the reduced expression of WISP-2 in colon tumors and cell lines suggests that it may function as 2 tumor suppressor. These results show that the WISP genes are aberrantly expressed in colon cancer and suggest that their altered expression may confer selective growth advantage to

Members of the Wnt signaling pathway have been implicated in the pathogenesis of colon cancer, breast cancer, and melanoma, including the tumor suppressor gene adenometous polyposis coli and B-catenin (39) Mutations in specific regions of either gene can cause the stabilization and accumulation of cytoplasmic B-catenin, which presumably contributes to human carcinogenesis through the activation of target genes such as the WISPs. Although the mechanism by which Writ-1 transforms cells and induces tumorigenesis is unknown, the Identification of WISPs as gones that may be regulated downstream of Wnt-1 in C57MG cells suggests they could be important mediators of Wnt-1 transformation. The amplificabon and altered expression patterns of the WISPs in human coion tumors may indicate an important role for these genes in tumor development.

We thank the DNA synthesis group for eligonucleotice synthesis. T. Baker for technical assistance, F. Dowd for resistion hybrid mapping. K. Willers and R. Nusso for the tee-ropessible CS7MG/Wnt-1 cells. V. Dixit for discussions, and D. Wood and A. Hruce for artwork.

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GENOMI METHODS

# Real Time Quantitative PCR

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We have developed a novel "real time" quantitative PCR method. The method measures PCR product accumulation through a dual-labeled fluorogenic probe (i.e., TogMan Probe). This method provides very accurate and reproducible quantitation of gene copies. Unlike other quantitative PCR methods, real-time PCR does not require post-PCR sample handling, preventing potential PCR product carry-over contamination and resulting in much faster and higher throughput assays. The real-time PCR method has a very large dynamic range of starting target molecule determination (at least live orders of magnitude). Real-time quantitative PCR is extremely accurate and less labor-intensive than current quantitative PCR methods.

Quantitative mudete acid sequence attalysis has had an important role in many fields of hiological research. Measurement of gene expression (RNA) has been used extensively in monitoring biological responses to various stimuli (l'an el al. 1994; Huang et al. 1995a,h; Prud'homme et al. 1995). Quantitation gene analysis (DNA) has been used to determine the gununu quantity of a particular gene, as in the case of the human HER2 gene, which is amplified in -30% of breast tumors (Slamon et al. 1987). Gene and genome quantitation (DNA and PNA) also have been used for analysis of human immunodeficiency virus (IIIV) buiden demonstrating changes in the levels of virus throughout the different phases of the disease (Connor et al. 1993; Plutak et al. 1993); Furtado et al. 1995).

Many methods have been described for the quantitative analysis of miciele acid sequences (both for RNA and DNA; Southern 1975; Sharp et al. 1980; Thomas 1980). Recently, PCIR has proven to be a powerful tool for quantitative nucleic acid analysis. PCR and reverse transcriptuse (RT)-PCR have permitted the analysis of minimal starting quantities of nucleic acid (as little as one cell equivalent). This has made possible many experiments that could not have been performed with traditional methods. Although PCR has provided a powerful tool, it is imperative

that it to used properly for quantitation (Rady-mackers 1995). Many early raports of quantitation of the PCR and RT-PCR described quantitation of the PCR product but did not measure the initial target sequence quantity. It is essential to design proper controls for the quantitation of the initial inter-sequences (Perre 1992; Clement) et al. 1003

Remarchers have developed several methods of quantitative PCR and RT-PCR. One approach measures PCR product quantity in the log phase of the reaction before the plateau (Kellogg et al. 1990; Pang et al. 1990). This method requires that each sample has equal input amounts of nucleic acid and that each sample under analysis amplifies with identical efficiency up to the point of quantitative analysis. A gene sequence (contalised in all samples of relatively constant quanlities, such as p-actin) can be used for sample umilification efficiency normalization. Using conventional methods of PCR detection and quantitation (gel electrophoresis or plate capture hybridization), it is extremely laborious to assure that all samples are analyzed during the log phase of the reaction (fur both the target gene and the normalization gone). Another method, quantitative competitive (QC)-PCR, has been developed and foused widery for PCR quantitation. QC-PCR relies on the inclusion of an internal control competitor in each reaction (Banker-Andre 1991; Matek et al. 1993a,b). The efficiency of each reaction is normalized to the Internal competitor. A known amount of internal competitor can be

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added to each sample. To obtain relative firstite totton, the unknown target PCR product is compared with the known competitor PCR product. Success of a quantitative competitive PCR assay relies on developing an internal control that amplifies with the same efficiency as the target molecule. The design of the compation and the validation of amplification efficiencies require a dedicated effort. However, because QC—PCR does not require that PCR products be analyzed during the log phase of the amplification, It is the eacher of the two methods to use.

Several detection systems are used for quanthative PCH and RICPCH analysis (1) agarase gels, (2) freorescent littlelling of PCR products and detection with insur-induced fluorescence using capillary electroplicresis (Fusco et al. 1995; WII-Homs et al. 1996) or acrylamide gels, and (3) place capture and sandwich probe hybridization (Muldar at al. 1994). Atthough these methods proved successful, each method requires post-PCR manipulations that add time to the analysis and may lead to Inhuratory contamination. The sample throughput of these methods is limited (with the exception of the plate capture approach), and, therefore, these methods are not well suited for uses demanding high sample throughput (i.e., screening of large numbers of libraniecules or analyxing samples for diagnostien or clinical trials).

Here we report the development of a novel ussay for quantitative DNA analysis. The array is based on the use of the 51 nucleuse assay first described by Holland et al. (1991). The method uses the 51 nuclease activity of Tray polymerate to cleave a nonextendible hybridization probe during the extension phase of PCR. The approach uses dual-labeled fluorogenic hybridization probes (Lee et al. 1983; Buscler et al. 1995; Livels et al., 1995a,b). One fluoreseent dye acres as a reporter (PAM (I.e., 6-carboxyfluoresectn)) and its emission spectra is quenched by the second flucrescent dye, TAYIRA (I.e., G-carlwixy-tetramethylrhodamine). The nuclease degradation of the hybridization probe releases the quenching of the PAM fluorescent emission, resulting in an increase in peak fluorescent emission at \$15 nm. The use of a sequence detector (ABI Prisin) allowe measurement of fluorescent appetra of all 96 wells of the thermal cyclet continuously during the PCR amplification. Therefore, the reactions are monitored in real time. The output data is described and quantitative unalysis of input larget INA sequences is discussed below.

#### RESLII.TS

## PCR Product Detection in Rual Time

The goal was to develop a high-throughput, senzitive, and accurate gene quantitation assay for use to monitoring lipid mudiated tharapoutic gene delivery. A plasmid uncoding human factor VIII geno requence, plisTM (see Methods), was used as a model therepeutic gene. The assay uses fluercheont Taquian methodology and an instrument capable of measuring fluorescence in real time (Ald Prism 7700 Sequence Descript). The Taymen reaction requires a hybridization probe isheled with two different fluorescent dyes. One dye is a reporter dye (HAM), the other is a quenching dye (TAMRA). When the probe is intact flucrescent energy transfer occurs and the reporter dye fluorescent emission is absorbed by the quenching dye (TAMRA). During the extension phase of the PCK cycle, the fluorescent hybridteation probe is cleaved by the 5'-3' nucleolytic activity of the DNA polymerase. On cleavage of the probe, the reporter dye emission is no longer transferred efficiently to the quenching dye, to sulting in an increase of the reporter dye fluores sont emission spectra. POR primare and probes were designed for the human factor VIII sequence and human p-actin gane (as described in Methods). Optimization reactions were performed to choose the appropriate probe and magnesium concentrations yielding the highest intensity of reporter fluorescent signal without encellicing specificity. The instrument once a charge-coupled device (i.e., CCD camera) for measuring the fluorescent emission spectra from 500 to 650 pm. Bach IYR tube was monitored sequentially for 25 msec with continuous munitoring throughout the amplification. Bach tube was re-examined every 8.5 see. Computer softwhre was designed to examine the fluorescent intensity of both the reporter dyn (FAM) and the quenching dyc (TAMIA). The Huorescent intensity of the quanching dys, TAMIU, changes very tittle over the course of the PCR amplification (data not shown). Therefore, the intensity of TAMBA due consistor serves as an Internal standard with which to normalise the reporter dye (FAM) emission variations. The software ealculules a value termed ARn (or ARQ) using the following equation: ARn = (Iln') (Iln'), where Rn4 .. emission intensity of reporter/emission intensity of quencher at any given time in a reaction tube, and Ru - emission intensitity of rePHONE No. : 318 472 8985

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porter/emission intensity of quencher measured prior to PCR amplification in that same reaction tube. For the purpose of quantitation, the last three data points (altris) conceted during the extension step for each PCR cycle were analyzed. The nucleolytic degradation of the nyundization probe occurs during the extension phase of PCR, and, therefore, reporter fluorescent emission increases during this time. The three data points were averaged for each PCR cycle and the mean value for each was plotted in an "amplification plot" shown in Figure 1A. The Alto mean value is plotted on the years, and time, represented by cycle number, is plotted on the years, and time, represented by cycle number, is plotted on the x-axis. During the early cycles of the FCR amplification, the Alto

value remains at base line. When sufficient hybridization probe has been cleaved by the Tan polymerase nuclease activity, the intensity of reporter fluorescent emission increases. Most PCR unplifications reach a plateau phase of reporter fluorescent emission if the reaction is carried and to high cycle muniscs. The emplification plot is examined early in the reaction, at a point that represents the log phase of predict accumulation. This is doing by assigning an arbitrary introsteoid that is laused on the variability of the base-time data. In Figure 1A, the threshold was set at its standard devolutions above the mean of base line emission calculated from typica 1 to 15. Once the threshold is chosen, the point at which

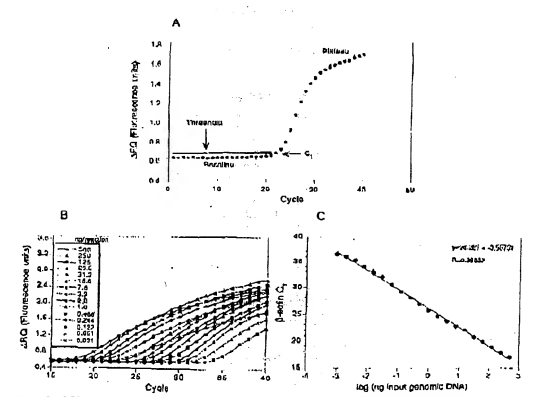


Figure 1. PCR product detection in real time. (A) The model 7700 sultiware will construct amplification plots from the extension phase fluorescent emission data collected during the PCR amplification. The standard deviation is determined from the data points collected from the base line of the amplification plot. C<sub>1</sub> values are calculated by determining the point at which the fluorescence exceeds a threshold limit (usually 10 times the standard deviation of the base line). (β) Overlay of amplification plots of serially (1:2) diluted human genomic DNA samples amplified with β-actin primers. (C) input DNA concentration of the samples plotted versus C<sub>7</sub>. All

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the amplification plot crosses the threshold wite fined as  $C_{\rm p}$ ,  $C_{\rm p}$  is reported as the cycle number of this point. As will be demonstrated, the  $C_{\rm p}$  value is predictive of the quantity of imput larget.

Cr Values Provide a Quantitative Measurement of Input Torget Sequences

Figure 18 shows amplification plots of 15 different PCR amplifications overlaid. The ampittantions were performed on a 1:2 serial dilutions of human genomic DNA. The amplified target was human B actin. The emplification plots shift to the right (to higher threshold cycles) as the input largel quantity is reduced. This is expected hucuited resections with fawer starting copins of the target molecule require greater amplification to degrade enough probe to attain the threshold fluorescence. An arbitrary threshold of 10 standard deviations above the base line was used to determine the C<sub>r</sub> values. Figure 1C represents the Cr values plotted versus the sample illuston value, Each dilution was amplified in triplicate PCR amplifications and plotted as mean values with error base representing one standard deviation. The Cryphes decrease linearly with increasing target quantity. Thus, Greature can be used as a quantitative measurement of the imput target number. It should be noted that the amplification plot for the 15.6-ng sample shown in Figure 1B does not reflect the same fluorescent rate of increase exhibited by most of the other samples. The 15.6-ng sample also actiteves endpoint piateau at a lower fluorescent value than would be expected based on the input DNA. This phenomenon has been observed occasionally with only samples (data not shown) and may be attributable to late cycle inhibition; this hypothesis is still under investigation. It is important to note that the flattened slope and carly plateau do not impact significantly the calculated Cq value as demonstrated by the fit on the line shown in Figure 1C. All triplicate amplifications resulted in very similar Ci values—the standard deviation did not exceed 0.5 for any dilution. This experiment contains a >100,000-fold range of Input target molecules. Using Co values for quantilation permits a much larger assay range than directly using total fluorescent emission intensity for quantitation. The linear range of fluorescent intensity measurement of the ABI Prism 7708 Sements over a very large range of relative starting target quantities.

#### Sample Preparation Validation

Several parameters influence the efficiency of PCR amplification: magnestum and sult concentrations, reaction conditions (i.e., time and tomparature), PCR target size and composition, printer sequences, and sample purity. All of the above factors are common to a single PCR assay, except sample to sample purity. In an effort to validate the method of sample preparation for the lactor VIII assay, PCR amplification reproducfollity and efficiency of 10 replicate sample preparations were examined. After genomic DNA was prepared from the 10 replicate samples, the DNA was quantitated by ultraviolet spectroscopy. Amplifications were performed analyzing p-actin gene content in 100 and 25 ng of total genomic DNA, Each PCR amplification was performed in triplicate. Comparison of C<sub>r</sub> values for each triplicate sample show minimal variation based on standard deviation and coefficient of variance (Pante 1). Therefore, each of the triplicate PCR amplifications was highly reproducible, demonstrating that real time PCR using this instrumentation introduces minimal variation into the quantitative l'Clt analysis. Comparison of the mean C, values of the 10 replicate sample preparations also showed minimal variability, indicating that each sample preparation yielded similar results for \$\beta\$-actin gene quantity. The highest Cr. difference between any of the samples was 0.85 and 0.71 for the 100 and 25 ng samples, resportively. Additionally, the amplification of each sample exhibited an equivalent rate of fluorescent emission intensity change per amount of DNA target analyzed as indicated by similar slopes derived from the sample dilutions (Fig. 2). Any sample containing an excess of a I'Ck inhibitor would exhibit a greater measured \$-actin Cr value for a given quantity of DNA. In addition, the inhibitor would be diluted along with the sample in the dilution analysis (Fig. 2), altering the expected C, value change. Each sample antplification yielded a similar result in the analysis, demonstrating that this method of sample proparation is highly reproducible with regard to sample purity.

Quantitative Analysis of a Plasmid After

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Sample		10	0 ng 🚟 🦈	. 47	25 ng				
	c <sub>t</sub>	mebn	standard deviation	cv	C,	mean	brabnata noitelvab	ÇV.	
1	18.24				20.48				
	18,23				20.55				
	76.33	14,27	0.06	0.32	20.5	20,51	0.03	0.17	
2	18.33				20.61				
	18.35	,			20.59				
	18.44	′ 18.37	0.06	0.32	20.41	20.54	0.11	0,51	
3	18.3				20.54				
	18.3				20,6				
	16.42	18.34	0.07	0.36	20.49	20.54	0. <b>0</b> 6	0.28	
4	18.15				20.48				
	18.23				20.44				
	18.32	18.23	30.0	0.45	20.38	20.43	0.05	0.26	
\$	18,4				20,68				
	18.38		:		20.87				
	18.46	18,42	0.04	0.23	20,63	20,71	0.13	0.61	
6	18.54				21.09				
	18.67				21.04				
	10	18.74	0,21	1.20	21.01	21.06	0,03	0.15	
7	18.28				20,67				
	18,36	•			20,73	•			
	18.52	18.39	0.12	0.66	<b>20</b> .65	20.68	0.04	0.2	
8	18.45				20,98				
	18.7				20.84	•	•		
	18,73	18,63	0.16	0.83	20.75	20.86	0.12	0.57	
9	18,18				20.46		•		
	18.34				20.54				
	18.36	18.29	0.1	0.55	20.48	20,51	למ,מ	0.32	
10	18.42				20.79				
	18,57				20.78				
	18,66	18.55	0.12	0.65	20.62	20.73	0.1	0.16	
Moan	(1 10)	18,12	0.17	0.90		20.66	0.19	0.94	

for containing a partial cDNA for human factor VIII, pERTM. A series of transfections was set up using a decreasing amount of the plasmid (40, 4, 0.5, and 0.1 µg). Twenty-four hours posttransfection, total DNA was purified from each flank of tells. p-Actin gene quantity was chosen as a value for normalization of genomic DNA concontration from each sample. In this experiment, Beactin sens content should remain constant relative to total genumic DNA. Figure 3 shows the result of the β-actin DNA measurement (100 mg total DNA determined by ultraviolet spectroscopy) of each sumple. Each sample was analyzed in idplicate and the mean p-actin Cr values of the triplicates were plotted (error bars represent men revealed neutrinoi The Monest difference

between any two sample moans was 0.95 C<sub>c</sub>. Ten hanograms of total DNA of each sample were also examined for fractin. The results again almoved that very similar amounts of genomic DNA were present; the modinum mean placin C<sub>t</sub> value difference was 1.0. As ligure 3 shows, the rate of placetic C<sub>t</sub> tilizings between the 100 and 10-ng samples was similar (slope values range between

3.56 and -3.45). This verifies again that the method of sample proparation yields samples of identical PCR integrity (i.e., no sample contained an oxcessive amount of a PCR inhibitor). However, these results indicate that each sample contained slight differences in the actual amount of genomic DNA analyzed, Determination of actual aground: DNA concentration was accomplished

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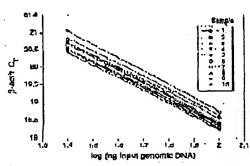


Figure 2 Sample preparation purity. The replicated samples shown in Table 1 wore also amplified in tripicate using 25 mg of each DNA sample. The figure shows the input DNA concentration (100 and 25 mg) vs. C. In the figure, the 100 and 25 mg points for each sample are connected by a fine.

by plotting the mean  $\beta$ -actin  $C_1$  value obtained for each 100-iig sample on a  $\beta$ -actin condard curve (shown in Fig. 4C). The eatial generale DNA concentration of each sample,  $\alpha$ , was obtained by extrapolation to the x-axis.

Figure 4A shows the measured (i.e., mornofmelined) quantities of factor VIII plasmid DNA (pretm) from each of the four transient cell transfections. Each reaction contained 100 ng of total sample DNA (as determined by UV spector copy). Each sample was analyzed in triplicate

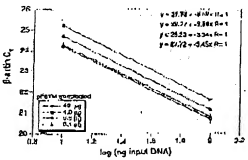


Figure 5. Analysis of transfected cell DNA quantity and purity. The DNA preparations of the four 293 cell transfections (40, 4, 0.5, and 0.1 µg of pF8TM) were analyzed for the 0-actin gene. 100 and 10 ng (determined by ultraviolet spectroscopy) of each sample were amplified in triplicate. For each amount of pF8TM that was transfected, the 6-actin Cryvalus are plotted versus the total input DNA

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PCR was pliffications. As shown, pl8TM purified strong the 293 cells decreases (mean C, values increases) with decreasing amounts of plasmid trumpleted. The mean C, values obtained for pp8TM in Figure 4A were plotted on a standard curve occupated of scribily diluted pp8TM, shown in Figure 4B. The quantity of post M, n, found in each of the four transfections was determined by extrapolation to the x axis of the standard curve in Figure 4B. These uncorrected values, n, for pP8TM were normalized to determine the actual amount of pP8TM found per 100 mg of genomic DNA by using the equation:

$$\frac{b \times 100 \text{ mg}}{a}$$
 = HCHIZI pPATM copies per 100 ng of genomic DNA

where a = actual generatic DNA in a sample and b = pF8TM copies from the standard curve. The normalized quantity of pF8TM per 100 ng of genomic ONA for each of the four transfections is snown in Figure 311. These results show that the quantity of factor VIII plasmid associated with the 250 cells, 24 in after transfection, decreases with decreasing plasmid contentation area in the transfection. The quantity of pF8TM associated with 253 cells, after transfection with 40 mg of plasmid, was 35 pg per 100 ng genomic DNA. This results in ~520 plasmid copies per cell.

#### DISCUSSION

We have described a new method for quantitating gene copy numbers using real-time analysis of PCR amplifications. Real-time PCR is compatible with either of the two PCR (ICT-PCR) approaches (1) quantitative competitive where an internal competitor for each target sequence is used for normalization (data not shown) or (2) quantitative comparative PCR using a normalization gene contained within the sample (i.e., β-action) or a "housekeeping" gene for RT-PCR. If equal amounts of nucleic acid are analyzed for each sample and if the amplification efficiency before quantitative analysis α identical for each sample, the internal control (normalization gene or competitor) should give equal agnals for all samples.

The real-time PCR method offers several advantages over the other two methods currently employed (see the introduction). First, the real-time PCR method is performed in a closed-tube system and requires no post-PCR manipulation

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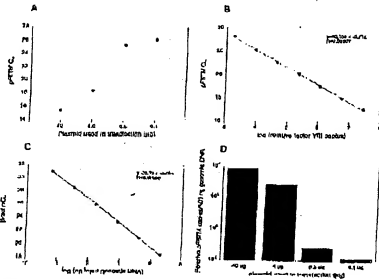


Figure 4. Quantitative analysis of pF8TM in transfected cells. (A) Amount of plasmid DNA used for the transfection plotted against the most C; value determined for pF8TM remaining 21 hr after transfection. (B,C) Standard curves of pE8TM and B-actin, respectively, pF8TM DNA (B) and generale DNA (C) were diluted socially 1:5 before amplification with the appropriate primers. The H-actin standard curve was used to normalize the results of A to 100 mg of genomic DNA. (D) The amount of pE8TM present per 100 mg of genomic DNA.

of sample. Therefore, the pertential for PCR confamination in the laboratory is reduced because amplifled products can be analyzed and disposed of without opening the reaction tobes. Second, this method supports the use of a normalization gona (i.e., H-actin) for quantitutive PCR or house keeping genes for quantitative RT-PCk controls. Analysis is performed in real time during the log phase of product accumulation. Analysis during lon phase permits many different genes (over a wide input target range) to be analyzed almultuneously, without concern of reaching reaction platom at different cycles. This will make multigone analysis assays much caster to develop, because individual internal competitors will not be needed for each gene under analysis. Third, sample throughput will increase dramatically with the new method because there is no post-PCR processing time. Additionally, wriking in a 96-well formst is highly compatible with automation technology.

The real-time PCR method is highly repreducible. Replicate amplifications can be analyzed

for each sample minimising potential error. The systems allows for a very large assay dynamic range (approaching 1,000,000-fold starting targot). Using a standard curve for the target of interest, relative copy number values can be determined for any unknown sample. Plugrescent threshold values, On correlate linearly with relative DNA copy numbers. Real time quantitative RT-PCR methodology (Gibson et al., this issue) has also been developed. Finally, real time quare thative I'CR mathodology can be used to develop high-throughput screening asserts for a variety of applications [quantitative gene unfamoion (1014 PCR), game copy amaya (Herž, IIIV, etc.), genertyping (knockout mouse analysis), and immune PCJY.

Real-time PCR may also be performed using Interculating dyes (Higachi et al. 1992) such us attaidium bromide. The fluorogenic probe method offers a major advantage over interculating dyes--greater specificity (i.e., primer dimers and nonspecific PCR products are not detected).

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## REAL TIME QUANTATIVE IVER

### METHODS

# Generation of a Plasmid Containing a Pertial CDNA for Human Factor VIII

Total RNA was harvated (RNArra D from To Teet, Inc., Irrendawood, TN) from colls transferred with a factor VIII repression vector, pCBS28c281 (Ration et al. 1986; Ourman et al. 1990). A factor VIII partial clina sequence was generated by RT PCB (Goneand) RC TTH RNA TCR RH (part N808-0179, T2 applied biosystems, roster City, Ca)] using the PCB genners Prior and Firev latinum acquirties are shown below). The amplicon was resimilated using modified Pifen and Pirev primers (appended with humilianed Hindill restricting sire sequences in the 5' quit and cloned into pclime 32 (Promaga Carp., Madison, WI). The resulting clone, pi6TM, was used for transfer transfection of 393 cells.

# Amplification of Target DNA and Detection of Amplicon Factor VIII Plasmed DNA

Amplification reactions (50 M) contained a DNA sample, TOX IXIL Buffer II (6 pl), 200 pm dATP, OCTU, dCTP, and 400 per dUTP, 4 mm MgCl., 1.25 Units Anspit Tag INA polymerate, O.S unit Amptrace urach neglyensylvac (UNC), 60 percels of each factor VIII printer, and 1.5 peticle of saids pain primes The reactions also consulted one of the following defection profits (100 nm epclis TRUTT(TAMPA) J' and A-netin probe 5' (FAM)ATGEXX: X(IAMIM)CCCCCATGCCATG1-31 where p indicates phosphorylation and X Indicmes a linker arm nuclearlite. Reaction tules were MicraAmp Optical Tules (part Aumber NEM 0933, Perion Blaser) that were frested (at Perion filmer) to present light from reflecting. Tube caps were similar to MicroAmp Claps but specially designed to prowent light scattering. All of the PCR communicables were sign which by Pli Applied Marystens (thater City, CA) except the factor VIII petruces, which were symbosized at Congn toch, Inc. (South San Francisco, CA), Probes were designed using the Oligo 4.0 software, following guidelings suggested in the Model 7700 Sequence Detector instituted munual, frieng, prope T\_ should be at least 5°C higher than the amealing temperature used during thermal cyching primers should not form stable duplexes with the

The thermal cycling conditions included 2 aim at 50°C and 10 min at 95°C. Thermal cycling proceeded with

reactions were performed in the Model 7200 Sequence Detector (IT: Applied Musystems), other cartains a Gene-Amp IICM System 1600. Reaction conditions were programmed up a Privat Macintosh 7100 (Apple Computor, Sama Clara, CA) United directly to the Model 9700 Sicquence Delector. Analysis of data was also preformed on the Medictock computer. Collection and analysis coffware was developed at 186 Applied Bloogiums.

## Transfection of Cells with Factor VIII Construct

Four T175 flasks of 293 criis GTCC CEL 1570. a human froit kidney suspension cell little, were grown to ROW, conthickey and transferred pittim, Cells were grown in the killawing media: SOM MAM'X F12 without G147, SOU law glucose Philheren's modified Fagie medium (DMRA) withoin glycline with sodium bicarbunate, 10% fetal bovine serum, 2 miss t-glutamine, and 106 penicillin-sureptomyelm. The media was changed 20 mile before the transfer sion, pHITM DNA amounts of 40, 4, 0.5, and 0.1 mg were idded to 1.6 ml of a solution containing 0.125 × CaCl2 and 1 x HIRPLS. The four mixtures were left at room lenepurature few 10 miles and then solded dropwine to the cally. The flasks were inculated at 27°C and FW CO2 for 24 hr, whished with PBS, and managemented in PBS. The toxics es saw AKCI been strongile of the block over cless thanks tracted immediately using the QIA sorp Blead Kit (Qlagon, Cliniamorth, (A). IDNA was chited into 200 pt of 20 min Tris-IICI at pH R.O.

### **ACKNOWLEDGMENTS**

We thank Generatech's DNA Synthesis Group for optimer synthesis and Generatech's Graphics Group for ostinance with the ligures

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Reactived hate 3, 1996; accepted in revised form July 29.

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methods. Peptides AENK or AEQK were dissolved in water, made isotonic with NaCl and diluted into RPMI growth medium. T-cell-proliferation assays were done essentially as described 10.11. Briefly, after antigen pulsing (30 µg ml-1 TTCF) with tetrapeptides (1-2 mg ml-1). PBMCs or EBV-B cells were washed in PBS and fixed for 45 s in 0.05% glutaraldehyde. Glycine was added to a final concentration of 0.1M and the cells were washed five times in RPMI 1640 medium containing 1% PCS before co-culture with T-cell clones in round-bottom 96-well microtitre plates. After 48 h, the cultures were pulsed with 1 µCi of 3M-thymidine and harvested for scintillation counting 16 h later. Predigestion of native TTCF was done by incubating 200 µg TTCF with 0.25 µg pig kidney legumain in 500 µl 50 mM citrate buffer, pH 5.5, for 1 h at 37 °C. Glycopoptide digestions. The peptides HIDNEEDI, HIDN(N-glucosamine) EEDI and HIDNESDI, which are based on the TTCF sequence, and QQQHLFGSNVTDCSGNFCLFR(KKK), which is based on human transferrin. were obtained by custom synthesis. The three C-terminal lysine residues were added to the natural sequence to aid solubility. The transferrin glycopeptide QQQHLFGSNVTDCSGNFCLFR was prepared by tryptic (Promega) digestion of 5 mg reduced, carboxy-methylated human transferrin followed by concanavalin A chromatography. Glycopeptides corresponding to residues 622-642 and 421-452 were isolated by reverse-phase HPLC and identified by mass spectrometry and N-terminal sequencing. The lyophilized transferrinderived peptides were redissolved in 50 mM sodium acetate, pH 5.5, 10 mM dithiothreitol, 20% methanol. Digestions were performed for 3 h at 30 °C with 5-50 mU ml-1 pig kidney legumain or B-cell AEP. Products were analysed by HPLC or MALDI-TOP mass spectrometry using a matrix of 10 mg ml-1 acyanocinnamic acid in 50% acetonitrile/0.1% TFA and a PerSeptive Biosystems cyanocinnamic acid in 50% acetoniumero. See the continued of the standard Elite STR mass spectrometer set to linear or reflector mode. Internal standard linear or reflector mode. The state of the s

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Acknowledgements. We drank M. Ferguson for helpful discussions and advice. E. Smythe and L. Crayson for advice and technical acciseance; B. Sprice, A. Knight and the BTS (Ninewalle Haspital) for help with blood monocyte preparation; and our colleagues for many helpful comments on the manuscripe. This

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work was supported by the Wellcome Trust and by an EMBO Long-term fellowship to B. M.

# Genomic amplification of a decoy receptor for Fas ligand in lung and colon cancer

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Fas ligand (Fast) is produced by activated T cells and natural killer cells and it induces apoptosis (programmed cell death) in target cells through the death receptor Fas/Apo1/CD95 (ref. 1). One important role of FasL and Fas is to mediate immunecytoroxic killing of cells that are potentially harmful to the organism, such as virus-infected or tumour cells'. Here we report the discovery of a soluble decoy receptor, termed decoy receptor 3 (DcR3), that binds to FasL and inhibits FasL-induced apoptosis. The DcR3 gene was amplified in about half of 35 primary lung and colon tumours studied, and DcR3 messenger RNA was expressed in malignant tissue. Thus, certain tumours may escape FasL-dependent immune-cytotoxic attack by expressing a decoy receptor that blocks Fasl.

By searching expressed sequence tag (EST) databases, we identified a set of related ESTs that showed homology to the tumour necrosis factor (TNF) receptor (TNFR) gene superfamily2. Using the overlapping sequence, we isolated a previously unknown fulllength complementary DNA from human fetal lung. We named the protein encoded by this cDNA decoy receptor 3 (DcR3). The cDNA encodes a 300-amino-acid polypeptide that resembles members of the TNFR family (Fig. 1a): the amino terminus contains a leader sequence, which is followed by four tandem cysteine-rich domains (CRDs). Like one other TNFR homologue, osteoprotegerin (OPG), DcR3 lacks an apparent transmembrane sequence, which indicates that it may be a secreted, rather than a membrane-associated. molecule. We expressed a recombinant, histidine-tagged form of DcR3 in mammalian cells; DcR3 was secreted into the cell culture medium, and migrated on polyacrylamide gels as a protein of relative molecular mass 35,000 (data not shown). DcR3 shares sequence identity in particular with OPG (31%) and TNFR2 (29%), and has relatively less homology with Fas (17%). All of the cysteines in the four CRDs of DcR3 and OPG are conserved; however, the carboxy-terminal portion of DcR3 is 101 residues shorter.

We analysed expression of DcR3 mRNA in human tissues by northern blotting (Fig. 1b). We detected a predominant 1.2-kilobase transcript in fetal lung, brain, and liver, and in adult spleen, coloa and lung. In addition, we observed relatively high DcR3 mRNA expression in the human colon carcinoma cell line SW480.

To investigate potential ligand interactions of DcR3, we generated a recombinant, Fc-tagged DcR3 protein. We tested binding of DcR3-Fc to human 293 cells transfected with individual TNIfamily ligands, which are expressed as type 2 transmembrane proteins (these transmembrane proteins have their N termini in the cytosol). DcR3-Fc showed a significant increase in binding to cells transfected with FasL' (Fig. 2a), but not to cells transfected with TNF', Apo2L/TRAIL', Apo3L/TWEAK', or OPGL/TRANCE/

RANKL<sup>16-12</sup> (data not shown). DcR3-Fc immunoprecipitated shed FasL from FasL-transfected 293 cells (Fig. 2b) and purified soluble FasL (Fig. 2c), as did the Fc-tagged ectodomain of Fas but not TNFR1. Gel-filtration chromatography showed that DcR3-Fc and soluble FasL formed a stable complex (Fig. 2d). Equilibrium analysis indicated that DcR3-Fc and Fas-Fc bound to soluble FasL with a comparable affinity ( $K_4 = 0.8 \pm 0.2$  and  $1.1 \pm 0.1$  nM, respectively; Fig. 2e), and that DcR3-Fc could block nearly all of the binding of soluble FasL to Fas-Fc (Fig. 2e, inset). Thus, DcR3 competes with Fas for binding to FasL.

To determine whether binding of DcR3 inhibits FasL activity, we tested the effect of DcR3-Fc on apoptosis induction by soluble FasL in Jurkat T leukaemia cells, which express Fas (Fig. 3a). DcR3-Fc and Fas-Fc blocked soluble-FasL-induced apoptosis in a similar dose-dependent manner, with half-maximal inhibition at -0.1 µg ml<sup>-1</sup>. Time-course analysis showed that the inhibition did not merely delay cell death, but rather persisted for at least 24 hours (Fig. 3b). We also tested the effect of DcR3-Fc on activation-induced cell death (AICD) of mature T lymphocytes, a FasL-dependent process'. Consistent with previous results', activation of interleukin-2-stimulated CD4-positive T cells with anti-CD3 antibody increased the level of apoptosis twofold, and Fas-Fc blocked this effect substantially (Fig. 3c); DcR3-Fc blocked the

induction of apoptosis to a similar extent. Thus, DcR3 binding blocks apoptosis induction by FasL.

FasL-induced apoptosis is important in elimination of virus-infected cells and cancer cells by natural killer cells and cytotoxic T lymphocytes; an alternative mechanism involves perforin and granzymes<sup>1,14-16</sup>. Peripheral blood natural killer cells triggered marked cell death in Jurkat T leukaemia cells (Fig. 3d); DcR3-Fc, and Fas-Fc each reduced killing of target cells from -65% to -30%, with half-maximal inhibition at -1 µg ml<sup>-1</sup>; the residual-killing was probably mediated by the perforin/granzyme pathway. Thus, DcR3 binding blocks FasL-dependent natural killer cell activity. Higher DcR3-Fc and Fas-Fc concentrations were required to block natural killer cell activity compared with those required to block soluble FasL activity, which is consistent with the greater potency of membrane-associated FasL compared with soluble FasL.

Given the role of immune cytotoxic cells in elimination of tumour cells and the fact that DoR3 can act as an inhibitor of Fast, we proposed that DoR3 expression might contribute to the ability of some tumours to escape immune-cytotoxic attack. As genomic amplification frequently contributes to tumorigenesis, we investigated whether the DoR3 gene is amplified in cancer. We analysed DoR3 gene-copy number by quantitative polymerase chain

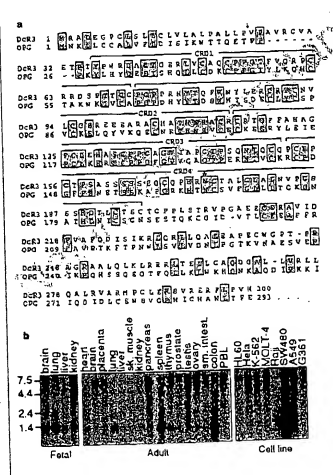


Figure 1 Primary structure and expression of human OcR3. a, Alignment of the amino-acid sequences of OcR3 and of osteoprotegerin (OPG); the C-terminal 101 residues of OPG are not shown. The putative signal cleavage site (arrow), the cysteme-rich domains (CR0 1-4), and the N-linked glycosylation site (asteriak) are shown. b, Expression of DcR3 mRNA. Northern hybridization analysis was done using the OcR3 cDNA as a probe and blots of poly(A)\* RNA (Clontech) from human fetal and adult tissuas or cancer cell fines. PBL, peripheral blood lymphocyte.

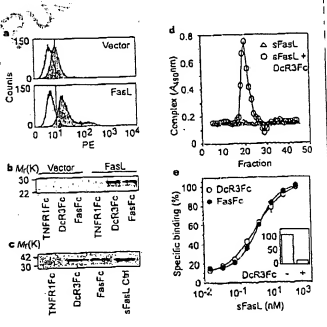


Figure 2 Interaction of DCR3 with Fast. a. 233 cella were transfected with pRK5 vector (top) or with pRK6 encoding full-length Fast. (bortom), incubated with DCR3-Fc (solld line, shaded area). TNFR1-Fc (dotted line) or buffer control (dashed line) (the dashed and dotted lines overlap), and analysed for binding by PACS. Statistical analysis showed a significant difference ( $t^2 < 0.001$ ) between the binding of DCR3-Fc to cells transfected with Fisal or pRK5. PE, phycoarythrilebelled cells, b, 293 cells were transfected as in a and metabolically labelled, and cell supermatants were immunoprecipitated with FC-tagged TNFR1, DCR3 or Fise. c. Purified soluble Fast. (sFast.) was immunoprecipitated with TNFR1-Fc, DCR3-Fc or Fas-Fc and visualized by immunoblot with anti-Fast entibody. aFast, was loaded directly for comparison in the right-hand lane. d. Flag-tagged aFast, was incubated with DCR3-Fc or with buffer and resolved by gel filtration; column tractions were analysed in an assay that detects complexes containing DcR2-Fc and sFast-Flag. e. Equilibrium binding of DcR3-Fc or Fas-Fc to sFast-Flag. Inset. competition of DcR3-Fc with Fas-Fc for binding to sFast-Flag.

reaction (PCR)<sup>16</sup> in genomic DNA from 35 primary lung and colon tumours, relative to pooled genomic DNA from peripheral blood leukocytes (PBLs) of 10 healthy donors. Eight of 18 lung tumours and 9 of 17 colon tumours showed DcR3 gene amplification, ranging from 2- to 18-fold (Fig. 4a, b). To confirm this result, we analysed the colon tumour DNAs with three more, independent sets of DcR3-based PCR primers and probes; we observed nearly the same amplification (data not shown).

We then analysed DcR3 mRNA expression in primary tumour tissue sections by in siru hybridization. We detected DcR3 expression in 6 out of 15 lung tumours. 2 out of 2 colon tumours, 2 out of 5 breast tumours, and 1 out of 1 gastric tumour (data not shown). A section through a squamous-cell carcinoma of the lung is shown in Fig. 4c. DcR3 mRNA was localized to infiltrating malignant epithelium, but was essentially absent from adjacent stroma, indicating tumour-specific expression. Although the individual tumour specimens that we analysed for mRNA expression and gene amplification were different, the in siru hybridization results are consistent with the finding that the DcR3 gene is amplified frequently in tumours. SW480 colon carcinoma cells, which showed abundant DcR3 mRNA expression (Fig. 1b), also had marked DcR3 gene amplification, as shown by quantitative PCR (fourfold) and by Southern blot hybridization (fivefold) (data not shown).

If DcR3 amplification in cancer is functionally relevant, then DcR3 should be amplified more than neighbouring genomic regions that are not important for tumour survival. To test this,

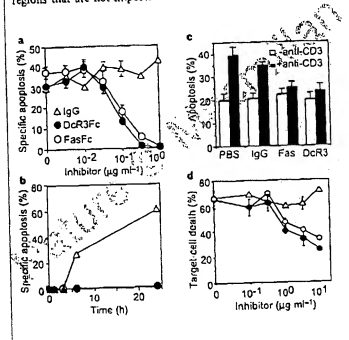


Figure 3 Inhibition of Fast, activity by DcR3, a, Human Jurkat T leukaemia cella were incubated with Flag-tagged soluble Fast (\$Fast; 5 ng ml<sup>-1</sup>) ollgomerized with anti-Flag antibody (0.1 µg ml<sup>-1</sup>) in the presence of the proposed inhibitors DcR3-Fc, Fas-Fc or human IgG1 and assayed for apoptosis (mean ± s.a.m. of triplicates), b, Jurket cells were incubated with 9Fast.—Flag plus anti-Flag antibody as in a, in presence of 1 µg ml<sup>-1</sup> DcR3-Fc (filled circles), Fas-Fc (open circles) or human IgG1 (triangles), and apoptosis was determined at the indicated time points. c, Feripheral blood T cells were stimulated with PHA and Interloukin-2, followed by control (white bers) or anti-CO3 entibody (filled bars), together with phosphate-buffered saline (PBS), human IgG1, Fas-Fc, or OcR3-Fc (10 µg ml<sup>-1</sup>). After 16 h, apoptosis of CO4\* cells was determined (mean ± s.e.m. of results from five donars). d, Paripheral blood natural killer cells were incubated with <sup>61</sup>Cr-Isbellod Jurkat cells in the presence of OcR3-Fc (filled circles), Fas-Fc (open circles) or human IgG1 (triangles), and targat-cell death was determined by release of CC (mean ± s.d. for two donors, each in triplicate).

we mapped the human DcR3 gene by radiation-hybrid analysis; DcR3 showed linkage to marker AFM218xe7 (T160), which maps to chromosome position 20q13. Next, we isolated from a bacterial artificial chromosome (BAC) library a human genomic clone that carries DcR3, and sequenced the ends of the clone's insert. We then determined, from the nine colon tumours that showed twofold or greater amplification of DcR3, the copy number of the DcR3flanking sequences (reverse and forward) from the BAC, and of seven genomic markers that span chromosome 20 (Fig. 4d) The DeR3-linked reverse marker showed an average amplification of roughly threefold, slightly less than the approximately fourfold amplification of DcR3; the other markers showed little or no amplification. These data indicate that Del3 may be at the 'epicentre' of a distal chromosome 20 region that is amplified in colon cancer, consistent with the possibility that DcR3 amplification promotes tumour survival.

Our results show that DcR3 binds specifically to FasL and inhibits FasL activity. We did not detect DcR3 binding to several other TNF-ligand-family members; however, this does not rule out the possibility that DcR3 interacts with other ligands, as do some other TNFR family members, including OPG<sup>217</sup>.

FasL is important in regulating the insmune response; however, little is known about how FasL function is controlled. One mechanism involves the molecule cFLIP, which modulates apoptosis signaling downstream of Fas<sup>30</sup>. A second mechanism involves proteolytic shedding of FasL from the cell surface<sup>17</sup>. DcR3 competes with Fas for

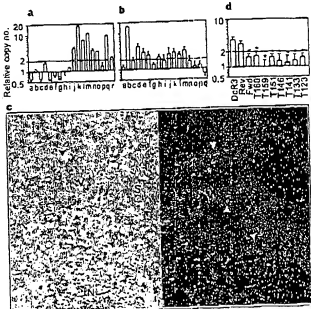


Figure 4 Genomic amplification of DcR3 in tumours, a, Lung cancers, comprising eight adenocarcinomas (c, d, f, g, h, ), k, r), sevan squamous-cell carcinomas (a, e, m, n, o, p, q), one non-small-cell carcinoma (b), one small-cell carcinoma (i), and one bronchiel adenocarcinoma (i). The data are means z s.d. of 2 experiments done in duplicate, b, Colon tumours, comprising 17 adenocarcinomes. Date are means = s.e.m. of five experiments done in duplicate. c. In situ hybridization analysis of DcR3 mRNA expression in a equamous-cell carcinoma of the lung. A representative bright-field image (left) and the corresponding dark-field Image (right) show DcR3 mRNA over Infiltrating meligriant apithelium (arrowheilds). Adjacent non-malignant stroma (S), blood veisel (V) and necrotic tumour tissue (N) are also shown, d, Average amplification of DcR3 compared with amplification of neighbouring genomic regions (reverse and forward, Rev and Fwd), the DcR3-linked marker T160, and other chromosoma-20 markers, in the nine colon tumours showing DcR3 amplification of twofold or more (b). Data are from two experiments done in duplicate. Asterisk indicates P < 0.01 for a Student's t-test comparing each marker with DcR3.

FasL binding; hence, it may represent a third mechanism of extracellular regulation of FasL activity. A decoy receptor that modulates the function of the cytokine interleukin-1 has been described14. In addition, two decoy receptors that belong to the TNFR family, DcR1 and DcR2, regulate the FasL-related apoptosisinducing molecule Apo2L12. Unlike DcR1 and DcR2, which are membranc-associated proteins, DcR3 is directly secreted into the extracellular space. One other secreted TNFR-family member is OPG', which shares greater sequence homology with DcR3 (31%) than do DcR1 (17%) or DcR2 (19%); OPG functions as a third decoy for Apo2L19. Thus, DcR3 and OPG define a new subset of TNFR-family members that function as secreted decoys to modulate ligands that induce apoptosis. Pox viruses produce soluble TNFR homologues that neutralize specific TNF-family ligands, thereby modulating the antiviral immune response2. Our results indicate that a similar mechanism, namely, production of a soluble decoy receptor for FasL, may contribute to immune evasion by certain tumours.

#### Mothods

Isolation of DcR3 cDNA. Several overlapping ESTs in GenBank (accession numbers AA025672, AA025673 and W67560) and in Lifeseq<sup>TM</sup> (Incyre Pharmaceuticals: accession numbers 1339238, 1533571, 1533650, 1542861, 1789372 and 2207027) showed similarity to members of the TNFR family. We screened human cDNA libraries by PCR with primers based on the region of EST consensus; fetal lung was positive for a product of the expected size. By hybridization to a PCR-generated probe based on the ESTs, one positive clones, (DNA30942) was identified. When scarching for potential alternatively spliced forms of DcR3 that might encode a transmembrane protein, we Isolated 500 more clones; the coding regions of these clones were identical in size to this of the initial clone (data not shown).

ectodomain of Fas or TNFR1, was fused to the hinge and Fc region of human

1gG1, expressed in insect SF9 cells or in human 293 cells, and purified as

described.

Fluorescence-activated cell sorting (FACS) analysis. We transfected 293 cells using calcium phosphate or Effectene (Qiagen) with pRKS vector or pRKS encoding full-length human Fash. (2 µg), together with pRKS encoding CrmA (2 µg) to prevent cell death. After 16 h, the cells were incubated with biotinylated DcR3-Fc or TNFR1-Fc and then with phycoerythrin-conjugated streptavidin (GibcoBRL), and were assayed by FACS. The data were analysed by Kolmogorov-Smirnov statistical analysis. There was some detectable staining of vector-transfected cells by DcR3-Fc; as these cells express little Fast (data not shown), it is possible that DcR3 recognized some other factor that is expressed constitutively on 293 cells.

Immunoprecipitation. Human 293 cells were transfected as above, and metabolically labelled with [35] cysteine and [35] methionine (0.5 mCi; Amersham). After 16 h of culture in the presence of z-VAD-fmk (10 µM), the medium was immunoprecipitated with DcR3-Fc, Fas-Fc or TNPR1-Fc (5 µg), followed by protein A-Sepharose (Repligen). The precipitates were resolved by SDS-PAGE and visualized on a phosphorimager (Fuji BAS2000). Alternatively, purified, Flag-tagged soluble Fast (1 µg) (Alexis) was incubated with each Fc-fusion protein (1 µg), precipitated with protein A-Sepharose, resolved by SDS-PAGE and visualized by immunoblotting with rabbit anti-Fast antibody (Oneogene Research).

Analysis of complex formation. Flag-tagged soluble FasL (25 µg) was incubated with buffer or with DcR3-Fc (40 µg) for 1.5 h at 24 °C. The reaction was loaded onto a Superdex 200 HR 10/30 column (Pharmacia) and developed with PBS; 0.6-ml fractions were collected. The presence of DcR3-Fc-FasL complex in each fraction was analysed by placing 100 µl aliquots into microtitre wells precoated with anti-human IgC (Boehringer) to capture DcR3-Fc, followed by detection with biotinylated anti-Flag antibody Bio M2 (Kodak) and streptavidin-horseradish peroxidase (Amersham). Calibration of the column indicated an apparent relative molecular mass of the complex of 420K (data not shown), which is consistent with a stoichiometry of two DcR3-Fc homodimers to two soluble FasL homotrimers.

Equilibrium binding analysis. Microtite wells were coated with anti-human

IgG, blocked with 2% BSA in PBS. DcR3-Fc or Fas-Fc was added, followed by serially diluted Flag-tagged soluble FasL. Bound ligand was detected with anti-Flag antibody as above. In the competition assay, Fas-Fc was immobilized as above, and the wells were blocked with excess IgG1 hefore addition of Flagtagged soluble FasL plus DcR3-Fc.

T-cell AICD. CD3\* lymphocytes were isolated from peripheral blood of individual donors using anti-CD3 magnetic beads (Miltenyi Biotech), stimulated with phytohaemagglutinin (PHA; 2 µg ml<sup>-1</sup>) for 24 h, and cultured in the presence of interleukin-2 (100 U ml<sup>-1</sup>) for 5 days. The cells were glated in wells coated with anti-CD3 antibody (Pharmingen) and analysed for anoptions 16 h later by FACS analysis of annexin-V-binding of CD4\* cells. The Natural killer cells were isolated from peripheral blood of individual donors using anti-CD56 imagnetic beads (Miltenyi Biotech), and incubated for 16 h with <sup>31</sup>Cr-loaded dickaticells at an effector-to-target ratio of 1:1 in the presence of DERJ-Pa; Fasi-Fc or human IgG1. Target-cell death was determined by release of <sup>51</sup>Cr in effector-target co-cultures relative to release of <sup>51</sup>Cr by degree thysis of equal numbers of Jurkat cells.

Gene-amplification analysis. Surgical specimens were provided by J. Kern (lung tumours) and P. Quirke (colon tumours). Genomic DNA was extracted (Qingen) and the concentration was determined using Hoethst dye 33258 intercalation fluorometry. Amplification was determined by quantitative PCR16 using a TaqMan instrument (ABI). The method was validated by comparison of PCR and Southern bybridization data for the Myc and HER-2 oncogenes (data not shown). Gene-specific primers and fluorogenic probes were designed on the basis of the sequence of DcR3 or of nearby regions identified on a BAC carrying the human DcRI gene: alternatively, primers and probes were based on Stanford Human Genome Center marker AFM218xe7 (T160), which is linked to DcR3 (likelihood score = 5.4), SHGC-36268 (T159), the nearest available marker which maps to ~500 kilobases from T160, and five extra markers that span chromosome 20. The DcR3-specific primer sequences were 5'-CTTCTTCGCGCACGCTG-3' and 5'-ATCACGCCGGCACCAG-3' and the fluorogenic probe sequence was 5'-(FAM-ACACGATGCGTGCTCCAAGCAG AAp-(TAMARA), where FAM is 5'-fluorescein phosphoramidite. Relative gene-copy numbers were derived using the formula 2(act), where act is the difference in amplification cycles required to detect DcR3 in peripheral blood lymphocyte DNA compared to test DNA.

## Received 24 September: accepted 6 November 1998.

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Acknowledgement. We also k C. Clark D. Pennica and V. Dixil for comments, and J. Kern and P. Quirke

Correspondence and requests for materials should be addressed to A.A. (e-mail: sa@gene.com). The GenBank accession number for the DCRJ cDNA sequence is AF104419.

# Crystal structure of the ATP-binding subunit of an ABC transporter

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ABC transporters (also known as traffic ATPases) form a large family of proteins responsible for the translocation of a variety of compounds across membranes of both prokaryotes and eukaryotes'. The recently completed Escherichia coli genome sequence revealed that the largest family of paralogous E coli proteins is composed of ABC transporters. Many enkiryotic proteins of medical significance belong to this family, such as the cystic fibrosis transmembrane conductance regulator (CFTR), the P-glycoprotein (or multidrug-resistance protein) and the heterodimeric transporter associated with antigen processing (Tap1-Tap2). Here we report the crystal structure at 1.5 A resolution of HisP, the ATP-binding subunit of the histidine permease, which is an ABC transporter from Salmonella typhimurium. We correlate the details of this structure with the biochemical, genetic and biophysical properties of the wild-type and several mutant HisP proteins. The structure provides a basis for understanding properties of ABC transporters and of defective CFTR proteins.

ABC transporters contain four structural domains: two nucleotide-binding domains (NBDs), which are highly conserved throughout the family, and two transmembrane domains! In prokaryotes these domains are often separate subunits which are assembled into a membrane-bound complex; in eukaryotes the domains are generally fused into a single polypeptide chain. The periplasmic histidine permease of S. typhimurium and E. coli 13-4 is a well-characterized ABC transporter that is a good model for this superfamily. It consists of a membrane-bound complex, HisQMP2, which comprises integral membrane subunits, HisQ and HisM, and two copies of HisP, the ATP-binding subunit. HisP, which has properties intermediate between those of integral and peripheral membrane proteins, is accessible from both sides of the membrane. presumably by its interaction with HisQ and HisM6. The two HisP subunits form a dimer, as shown by their cooperativity in ATP hydrolysis, the requirement for both subunits to be present for activity, and the formation of a HisP dimer upon chemical crosslinking. Soluble HisP also forms a dimer'. HisP has been purified and characterized in an active soluble form' which can be reconstituted into a fully active membrane-bound complex.

The overall shape of the crystal structure of the HisP monomer is that of an 'L' with two thick arms (arm I and arm II); the ATP-binding pocket is near the end of arm I (Fig. 1). A six-stranded  $\beta$ -sheet ( $\beta$ 3 and  $\beta$ 8- $\beta$ 12) spans both arms of the L, with a domain of a  $\alpha$ -plus  $\beta$ -type structure ( $\beta$ 1,  $\beta$ 2,  $\beta$ 4- $\beta$ 7,  $\alpha$ 1 and  $\alpha$ 2) on one side (within arm I) and a domain of mostly  $\alpha$ -helices ( $\alpha$ 3- $\alpha$ 9) on the

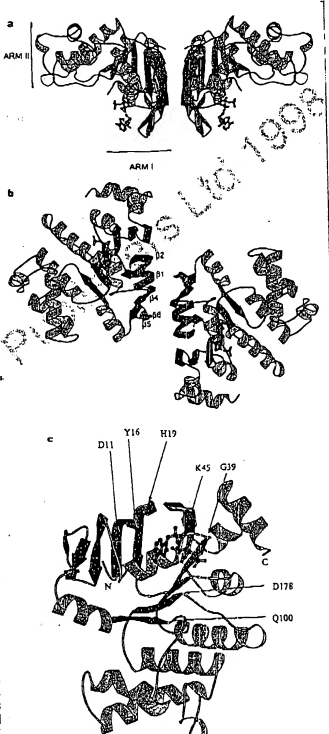


Figure 1 Crystal structure of HisP. 2 View of the dimer along an axis perpendicular to its two-fold exis. The top and bottom of the dimer are suggested to face towards the periplasmic and cymplasmic sides, respectively (see text). The thickness of arm flis about 25 Å, comparable to that of membrane, a-Halicus are shown in orange and p-sheete in green, b. View along the two-fold axis of the HisP dimer, showing the relative displacement of the monomers not apparent in a. The p-strande at the dimer interface are labelled o. View of one monomer from the bottom of erm I, as shown in a, towards erm II, showing the ATP-binding pocket, a-c. The protein and the bound ATP are in 'hibbon' and 'ball-and-stick' representations, respectively. Key residues discussed in the text are indicated in c. These figures were prepared with MOLSCRIPT<sup>24</sup>. N, emino terminus; C. C terminus.

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Int. J. Cancer: 78, 661-666 (1998) © 1998 Wiley-Liss, Inc.



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# NOVEL APPROACH TO QUANTITATIVE POLYMERASE CHAIN REACTION USING REAL-TIME DETECTION: APPLICATION TO THE DETECTION OF GENE AMPLIFICATION IN BREAST CANCER

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Gene amplification is a common event in the progression of human cancers, and amplified oncogenes have been shown to have diagnostic, prognostic and therapeutic relevance. A kinetic quantitative polymerase-chain-reaction (PCR) method, based on fluorescent TaqMan methodology and a new instrument (ABI Prism 7700 Sequence Detection System) capable of measuring fluorescence in real-time, was used to quantify gene amplification in tumor DNA. Reactions are characterized by the point during cycling when PCR amplification is still in the exponential phase, rather than the amount of PCR product accumulated after a fixed number of cycles. None of the reaction components is limited during the exponential phase, meaning that values are highly reproducible in reactions starting with the same copy number. This greatly improves the precision of DNA quantification. Moreover, real-time PCR does not require post-PCR sample handling, thereby preventing potential PCR-product carry-over contamination; it possesses a wide dynamic range of quantification and results in much faster and higher sample throughout. The real-time PCR method, was used to develop and validate a simple and rapid assay for the detection and quantification of the 3 most frequently amplified genes (myc, cond) and erbB2) in breast tumors. Extra copies of myc, cend1 and erbB2 were observed in 10, 23 and 15%, respectively, of 108 breasttumor DNA; the largest observed numbers of gene copies were 4.6, 18.6 and 15.1, respectively. These results correlated well with those of Southern blotting. The use of this new semi-automated technique will make molecular analysis of human cancers simpler and more reliable, and should find broad applications in clinical and research settings. Int. L Cancer 78:661-666, 1998. o 1998 Wiley-Liss, Inc.

Gene amplification plays an important role in the pathogenesis of various solid numors, including breast cancer, probably because over-expression of the amplified target genes confers a selective advantage. The first technique used to detect genomic amplification was cytogenetic analysis. Amplification of several chromosome regions, visualized either as extrachromosomal double minutes (dmins) or as integrated homogeneously staining regions (HSRs), are among the main visible cytogenetic abnormalities in breast tumors. Other techniques such as comparative genomic hybridization (CGH) (Kallioniemi et al., 1994) have also been used in broad searches for regions of increased DNA copy numbers in tumor cells, and have revealed some 20 amplified chromosome regions in breast tumors. Positional cloning efforts are underway to identify the critical gene(s) in each amplified region. To date, genes known to be amplified frequently in breast cancers include myc (8q24), cond1 (11q13), and erbB2 (17q12-q21) (for review, see Bieche and Lidereau, 1995).

Amplification of the myc. ccndl, and erbB2 proto-oncogenes should have clinical relevance in breast cancer, since independent studies have shown that these alterations can be used to identify sub-populations with a worse prognosis (Berns et al., 1992; Schuuring et al., 1992; Mamon et al., 1987). Muss et al. (1994) suggested that these gene alterations may also be useful for the prediction and assessment of the efficacy of adjuvant chemotherapy and hormone therapy.

However, published results diverge both in terms of the frequency of these alterations and their clinical value. For instance, over 500 studies in 10 years have failed to resolve the controversy

surrounding the link suggested by Slamon et al. (1987) between erbB2 amplification and disease progression. These discrepancies are partly due to the clinical, histological and ethnic heterogeneity of breast cancer, but technical considerations are also probably involved.

Specific genes (DNA) were initially quantified in tumor cells by means of blorting procedures such as Southern and slot blotting. These batch techniques require large amounts of DNA (5-10 µg/reaction) to yield reliable quantitative results. Furthermore, meticulous care is required at all stages of the procedures to generate blots of sufficient quality for reliable dosage analysis. Recently, PCR has proven to be a powerful tool for quantitative DNA analysis, especially with minimal starting quantities of tumor samples (small, early-stage tumors and formalin-fixed, paraffinembedded tissues).

Quantitative PCR can be performed by evaluating the amount of product either after a given number of cycles (end-point quantitative PCR) or after a varying number of cycles during the exponential phase (kinetic quantitative PCR). In the first case, an internal standard distinct from the target molecule is required to ascertain PCR efficiency. The method is relatively easy but implies generating, quantifying and storing an internal standard for each gene studied. Nevertheless, it is the most frequently applied method to date.

One of the major advantages of the kinetic method is its rapidity in quantifying a new gene, since no internal standard is required (an external standard curve is sufficient). Moreover, the kinetic method has a wide dynamic range (at least 5 orders of magnitude), giving an accurate value for samples differing in their copy number. Unfortunately, the method is cumbersome and has therefore been rarely used. It involves aliquot sampling of each assay mix at regular intervals and quantifying, for each aliquot, the amplification product. Interest in the kinetic method has been stimulated by a novel approach using fluorescent TaqMun methodology and a new instrument (ABI Prism 7700 Sequence Detection System) capable of measuring fluorescence in real time (Gibson et al., 1996; Heid et al., 1996). The TaqMan reaction is based on the 5' nuclease assay first described by Holland et al. (1991). The latter uses the S nuclease activity of Tsq polymerase to cleave a specific fluorogenic oligonucleotide probe during the extension phase of PCR. The approach uses dual-labeled fluorogenic hybridization probes (Lee et al., 1993). One fluorescent dye, co-valently linked to the 5' and of the oligonucleotide, serves as a reporter [FAM (i.e., 6-carboxyfluorescein)] and its emission spectrum is quenched by a second fluorescent dyc, TAMRA (i.e., 6-carboxy-tetramethyl-rhodamino) attached to the 3' end. During the extension phase of the PCR

Grant sponsors: Association Pour la Recherche sur le Cancer and Ministère de l'Enscignement Supérieur et de la Recherche.

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cycle, the fluorescent hybridization probe is hydrolyzed by the 5'-3' nucleolytic activity of DNA polymerase. Nuclease degradation of the probe releases the quenching of FAM fluorescence emission, resulting in an increase in peak fluorescence emission. The fluorescence signal is normalized by dividing the emission intensity of the reporter dye (FAM) by the emission intensity of a reference dye (i.e., ROX, 6-carboxy-X-rhodamine) included in TaqMan buffer, to obtain a ratio defined as the Rn (normalized reporter) for a given reaction tube. The use of a sequence detector enables the fluorescence spectra of all 96 wells of the thermal cycler to be measured continuously during PCR amplification.

The real-time PCR method offers several advantages over other current quantitative PCR methods (Celi et al., 1994): (i) the probe-based homogeneous assay provides a real-time method for detecting only specific amplification products, since specific hybridation of both the primers and the probe is necessary to generate a signal; (ii) the C, (threshold cycle) value used for quantification is measured when PCR amplification is still in the log phase of PCR product accumulation. This is the main reason why C<sub>i</sub> is a more reliable measure of the starting copy number than are end-point measurements, in which a slight difference in a limiting component can have a drastic effect on the amount of product; (iii) use of C, values gives a wider dynamic range (at least 5 orders of magnitude), reducing the need for serial dilution; (iv) The real-time PCR method is run in a closed-rube system and requires no post-PCR sample handling, thus avoiding potential contamination; (v) the system is highly automated, since the instrument continuously measures fluorescence in all 96 wells of the thermal cycler during PCR amplification and the corresponding software processes, and analyzes the fluorescence data; (vi) the assay is rapid, as results are available just one minute after thormal cycling is complete; (vii) the sample throughput of the method is high, since 96 reactions can be analyzed in 2 hr.

Here, we applied this semi-automated procedure to determine the copy numbers of the 3 most frequently amplified genes in breast tumors (myc, cend1 and erbB2), as well as 2 genes (alb and app) located in a chromosome region in which no genetic changes have been observed in breast tumors. The results for 108 breast tumors were compared with previous Southern-blot data for the same samples.

## MATERIAL AND METHODS

Tumor and blood samples

Samples were obtained from 108 primary breast tumors removed surgically from patients at the Centre Rene Huguenin; none of the patients had undergone radiotherapy or chemotherapy. Immediately after surgery, the itumor samples were placed in liquid nitrogen until extraction of high-molecular-weight DNA. Patients were included in this study if the tumor sample used for DNA preparation contained more than 60% of tumor cells (histological analysis). A blood sample was also taken from 18 of the same natients.

DNA was extracted from tumor tissue and blood loukocytes according to standard methods.

## Real-time PCR

Theoretical basis. Reactions are characterized by the point during cycling when amplification of the PCR product is first detected, rather than by the amount of PCR product accumulated after a fixed number of cycles. The higher the starting copy number of the genomic DNA target, the earlier a significant increase in fluorescence is observed. The parameter C<sub>1</sub> (threshold cycle) is defined as the fractional cycle number at which the fluorescence generated by cleavage of the probe passes a fixed threshold above baseline. The target gene copy number in unknown samples is quantified by measuring C<sub>1</sub> and by using a standard curve to determine the starting copy number. The precise amount of genomic DNA (based on optical density) and its quality (i.e., lack

of extensive degradation) are both difficult to assess. We therefore also quantified a control gene (alb) mapping to chromosome region 4q11-q13, in which no genetic alterations have been found in breast-numor DNA by means of CGH (Kallioniemi et al., 1994).

Thus, the ratio of the copy number of the target gene to the copy number of the alb gene normalizes the amount and quality of genomic DNA. The ratio defining the level of amplification is termed "N", and is determined as follows:

N = copy number of target gene (app. myc. candl. erbB2)
copy number of reference gene (alb)

Primers, probes, reference human genomic DNA and PCR consumables. Primers and probes were chosen with the assistance of the computer programs Oligo 4.0 (National Biosciences, Plymouth, MN), EuGene (Daniben Systems, Cincinnati, OH) and Primer Express (Perkin-Elmer Applied Biosystems, Foster City, CA).

Primers were purchased from DNAgency (Malvern, PA) and probes from Perkin-Elmer Applied Biosystems.

Nucleotide sequences for the oligonucleotide hybridization probes and primers are available on request.

The TaqMan PCR Core reagent kit, MicroAmp optical tubes, and MicroAmp caps were from Perkin-Elmer Applied Biosystems.

Standard-curve construction. The kinetic method requires a standard curve. The latter was constructed with serial dilutions of specific PCR products, according to Piatak et al. (1993). In practice, each specific PCR product was obtained by amplifying 20 ng of a standard human genomic DNA (Boehringer, Mannheim, Germany) with the same primer pairs as thuse used later for real-time quantitative PCR. The 5 PCR products were purified using MicroSpin S-400 HR columns (Pharmacia, Uppsala, Sweden) electrophorezed through an acrylamide gel and stained with ethidium bromide to check their quality. The PCR products were then quantified spectrophotometrically and prooled, and serially diluted 10-fold in mouse genomic DNA (Clontech, Palo Alto, CA) at a constant concentration of 2 ng/ul. The standard curve used for real-time quantitative PCR was based on serial dilutions of the pool of PCR products ranging from  $10^{-7}$  (10° copies of each gene) to 10-10 (102 copies). This series of diluted PCR products was aliquoted and stored at -80°C until use.

The standard curve was validated by analyzing 2 known quantities of calibrator human genomic DNA (20 ng and 50 ng).

PCR amplification. Amplification mixes (50 µl) contained the sample DNA (around 20 ng, around 6600 copies of disomic gence), 10× TaqMan buffer (5 µl), 200 µM dATP, dCTP, dGTP, and 400 µM dUTP, 5 mM MgCl<sub>2</sub>, 1.25 units of AmpliTaq Gold, 0.5 units of Amplerase uracil N-glycosylase (UNG), 200 nM each primer and 100 nM probe. The thermal cycling conditions comprised 2 min at 50°C and 10 min at 95°C. Thermal cycling consisted of 40 cycles at 95°C for 15 s and 65°C for 1 min. Each assay included: a standard curve (from 10° to 10² copies) in duplicate, a no-template control, 20 ng and 50 ng of calibrator human genomic DNA (Boehringer) in miplicate, and about 20 ng of unknown genomic DNA in triplicate (26 samples can thus be analyzed on a 96-well microplate). All samples with a coefficient of variation (CV) higher than 10% were

All reactions were performed in the ABI Prism 7700 Sequence Detection System (Perkin-Elmer Applied Biosystems), which detects the signal from the fluorogenic probe during PCR.

Equipment for real-time detection. The 7700 system has a built-in thermal cycler and a laser directed via fiber optical cables to each of the 96 sample wells. A charge-coupled-device (CDD) camera collects the emission from each sample and the data are analyzed automatically. The software accompanying the 7700 system calculates C<sub>1</sub> and determines the starting copy number in the samples.

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Determination of gene amplification. Gene amplification was calculated as described above. Only samples with an N value higher than 2 were considered to be amplified.

### RESULTS

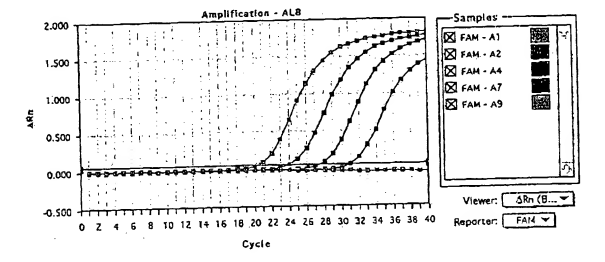
To validate the method, real-time PCR was performed on genomic DNA extracted from 108 primary breast tumors, and 18 normal leukocyte DNA samples from some of the same patients. The target genes were the *mye*, *cend1* and *erbB2* proto-oncogenes, and the β-amyloid precursor protein gene (*app*), which maps to a chromosome region (21q21.2) in which no genetic alterations have been found in breast tumors (Kallioniemi *et al.*, 1994). The reference disonic gene was the albumin gene (*alb.* chromosome 4q11-q13).

Validation of the standard curve and dynamic range of real-time PCR

The standard curve was constructed from PCR products scrially diluted in genomic mouse DNA at a constant concentration of 2 ng/µl. It should be noted that the 5 primer pairs chosen to analyze the 5 target genes do not amplify genomic mouse DNA (data not shown). Figure 1 shows the real-time PCR standard curve for the alb gene. The dynamic range was wide (at least 4 orders of magnitude), with samples containing as few as 10<sup>2</sup> copies or as many as 10<sup>5</sup> copies.

Copy-number ratio of the 2 reference yenes (app and alb)

The app to alb copy-number ratio was determined in 18 normal leukocyte DNA samples and all 108 primary breast-numor DNA



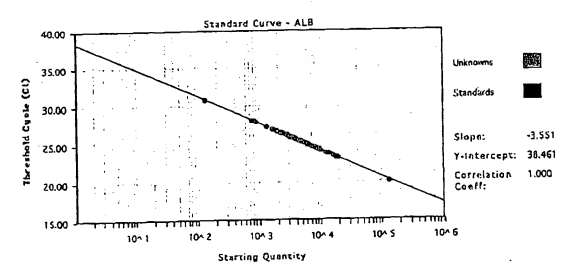


FIGURE 1 - Albumin (alb) gene dosage by real-time PCR. Top: Amplification plots for reactions with starting alb gene copy number ranging from 10<sup>5</sup> (A9), 10<sup>4</sup> (A7), 10<sup>3</sup> (A4) to 10<sup>5</sup> (A2) and a no-template control (A1). Cycle number is plotted vs. change in normalized reporter signal (ARn). For each reaction tube, the fluorescence signal of the reporter dye (FAM) is divided by the fluorescence signal of the passive reference dye (ROX), to obtain a ratio defined as the normalized reporter signal (Rn). ARn represents the normalized reporter signal (Rn) minus the baseline signal established in the first 15 PCR cycles. ARn increases during PCR as alb PCR product copy number increases until the reaction reactes a signal established in the first 15 PCR cycles. ARn increases during PCR as alb PCR product copy number increases until the reaction reactes plateau. C<sub>1</sub> (threshold cycle) represents the fractional cycle number at which a significant increase in Rn above a baseline signal (horizontal black plateau. C<sub>2</sub> (threshold cycle) represents the fractional cycle number at which a significant increase in Rn above a baseline signal (horizontal black plateau. C<sub>3</sub> (threshold cycle) represents the fractional cycle number at sandard sample, but the data for only one are shown here. Bottom: Item fractions are plotting log starting copy number vs. C<sub>3</sub> (threshold cycle). The black dots represent the data for standard samples plotted in triplicate. The standard curve shows 4 orders of linear dynamic tange.

samples. We selected these 2 genes because they are located in 2 chromosome regions (app. 21q21.2; alb. 4q11-q13) in which no obvious genetic changes (including gains or losses) have been observed in breast cancers (Kallioniemi et al., 1994). The ratio for the 18 normal leukocyte DNA samples fell between 0.7 and 1.3 (mean  $1.02 \pm 0.21$ ), and was similar for the 108 primary breast-tumor DNA samples (0.6 to 1.6, mean  $1.06 \pm 0.25$ ), confirming that alb and app are appropriate reference disomic genes for breast-tumor DNA. The low range of the ratios also confirmed that the nucleotide sequences chosen for the primers and probes were not polymorphic, as mismatches of their primers or probes with the subject's DNA would have resulted in differential amplification.

myc, condl and crbB2 gene dase in normal leukocyte DNA

To determine the cut-off point for gene amplification in breast-cancer tissue, 18 normal leukocyte DNA samples were tested for the gene dose (N), calculated as described in "Material and Methods". The N value of these samples ranged from 0.5 to 1.3 (mean 0.84 ± 0.22) for myc. 0.7 to 1.6 (mean 1.06 ± 0.23) for cend1 and 0.6 to 1.3 (mean 0.91 ± 0.19) for erbB2. Since N values for myc. cend1 and erbB2 in normal leukocyte DNA consistently fell between 0.5 and 1.6, values of 2 or more were considered to represent gene amplification in tumor DNA.

myc. ccndl and etbB2 gene dose in breast-tumor DNA

myc, cond1 and erbB2 genc copy numbers in the 108 primary breast tumors are reported in Table I. Extra copies of cond1 were more frequent (23%, 25/108) than extra copies of erbB2 (15%, 16/108) and myc (10%, 11/108), and ranged from 2 to 18.6 for cond1, 2 to 15.1 for erbB2, and only 2 to 4.6 for the myc gene. Figure 2 and Table II represent tumors in which the cond1 gene was amplified 16-fold (T145), 6-fold (T133) and non-amplified (T118). The 3 genes were never found to be co-amplified in the same tumor. erbB2 and cond1 were co-amplified in only 3 cases, myc and cond1 in 2 cases and myc and erbB2 in 1 case. This favors the hypothesis that gene amplifications are independent events in breast cancer. Interestingly, 5 tumors showed a decrease of at least 50% in the erbB2 copy number (N < 0.5), suggesting that they bore deletions of the 17q21 region (the site of erbB2). No such decrease in copy number was observed with the other 2 proto-oncogenes.

Camparison of gene dose determined by real-time quantitative PCR and Southern-blot analysis

Southern-blot analysis of myc, cend1 and erbB2 amplifications had previously been done on the same 108 primary breast tumors. A perfect correlation between the results of real-time PCR and Southern blot was obtained for tumors with high copy numbers  $(N \ge 5)$ . However, there were cases (1 myc, 6 cend1) and 4 erbB2 in which real-time PCR showed gene amplification whereas Southern-blot did not, but these were mainly cases with low extra copy numbers (N from 2 to 2.9).

## DISCUSSION

The clinical applications of gene amplification assays are currently limited, but would certainly increase if a simple, standardized and rapid method were perfected. Gene amplification status has been studied mainly by means of Southern blotting, but this method is not sensitive enough to detect low-level gene amplification nor accurate enough to quantify the full range of amplification values. Southern blotting is also time-consuming, uses radioactive

TABLE I - DISTRIBUTION OF AMPLIFICATION LEVEL (N) FOR IMPC. cends and erobs genes in 108 Human Breast TUMORS

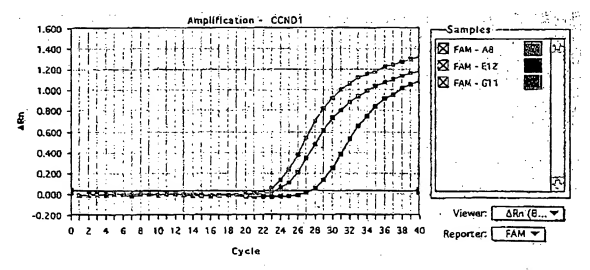
_	Amplification level (N)					
Gene	<0.5	0.5-1.9	2_4.9	≥5		
myc	0	97 (89.8%)	11 (10.2%)	0		
ccnd1	0	83 (76.9%)	17 (15.7%)	8 (7.4%)		
erbBZ	5 (4.6%)	87 (30.6%)	8 (7.4%)	8 (7.4%)		

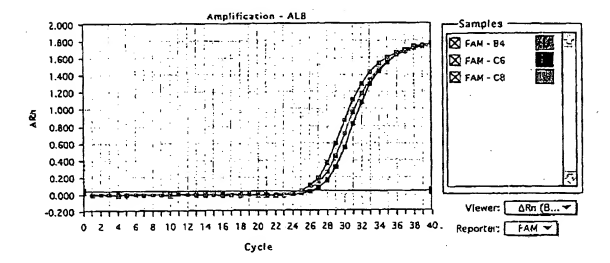
reagents and requires relatively large amounts of high-quality genomic DNA, which means it cannot be used routinely in many laboratories. An amplification step is therefore required to determine the copy number of a given target gene from minimal quantities of tumor DNA (small early-stage tumors, cytopuncture specimens or formalin-fixed, paraffin-embedded tissues).

In this study, we validated a PCR method developed for the quantification of gene over-representation in tumors. The method, based on real-time analysis of PCR amplification, has several advantages over other PCR-based quantitative assays such as competitive quantitative PCR (Celi et al., 1994). First, the real-time PCR method is performed in a closed-tube system, avoiding the risk of contamination by amplified products. Re-amplification of carryover PCR products in subsequent experiments can also be prevented by using the enzyme uracil N-glycosylase (UNG) (Longo et al., 1990). The second advantage is the simplicity and rapidity of sample analysis, since no post-PCR manipulations are required. Our results show that the automated method is reliable. We found it possible to determine, in triplicate, the number of copies of a rarget gene in more than 100 rumors per day. Third, the system has a linear dynamic range of at least 4 orders of magnitude, meaning that samples do not have to contain equal starting amounts of DNA. This technique should therefore be suitable for analyzing formalin-fixed, paraffin-embedded tissues. Fourth, and above all real-time PCR makes DNA quantification much more precise and reproducible, since it is based on C, values rather than end-point measurement of the amount of accumulated PCR product. Indeed, the ABI Prism 7700 Sequence Detection System enables C, to be calculated when FCR amplification is still in the exponential phase and when none of the reaction components is rate-limiting. The within-run CV of the C, value for calibrator human DNA (5 replicates) was always below 5%, and the between-assay precision in 5 different runs was always below 10% (data not shown). In addition, the use of a standard curve is not absolutely necessary, since the copy number can be determined simply by comparing the Ciratio of the target gene with that of reference genes. The results obtained by the 2 methods (with and without a standard curve) are similar in our experiments (data not shown). Moreover, unlike competitive quantitative PCR, real-time PCR does not require an internal control (the design and storage of internal controls and the validation of their amplification efficiency is laborious).

The only potential disavantage of real-time PCR, like all other PCR-based methods and solid-matrix blotting techniques (Southem blots and dot blots) is that is cannot avoid dilution artifacts inherent in the extraction of DNA from temor cells contained in heterogeneous tissue specimens. Only FISH and immunohistochemistry can measure alterations on a cell-by-cell basis (Pauletti et al., 1996; Slamon et al., 1989). However, FISH requires expensive equipment and trained personnel and is also time-consuming. Moreover, FISH does not assess gene expression and therefore cannot detect cases in which the genc product is over-expressed in the absence of gene amplification, which will be possible in the future by real-time quantitative RT-PCR. Immunohistochemistry is subject to considerable variations in the hands of different teams, owing to alterations of target proteins during the procedure, the different primary antibodies and fixation methods used and the criteria used to define positive staining.

The results of this study are in agreement with those reported in the literature. (i) Chromosome regions 4q11-q13 and 21q21.2 (which bear alh and app, respectively) showed no genetic alterations in the breast-cancer samples studied here, in keeping with the results of CGH (Kallioniemi et al., 1994). (ii) We found that amplifications of these 3 oncogenes were independent events, as reported by other teams (Berns et al., 1992; Borg et al., 1992). (iii) The frequency and degree of myc amplification in our breast tumor DNA series were lower than those of cend1 and erbB2 amplification, confirming the findings of Borg et al. (1992) and Courjal et al. (1997). (iv) The maxima of cend1 and erbB2 over-representation were 18-fold and 15-fold, also in keeping with earlier results (about





	(	CCND1	ALB		
Tumor	$C_{t}$	opy number	Ct C	Copy number	
T118	27.3	4605	26.5	4365	
<b>國 T133</b>	23.2	61659	25.2	10092	
<b>■ T145</b>	22.1	125892	25.6	7762	

FIGURE 2 - cond1 and alb gene dosage by real-time PCR in 3 breast tumor samples: T118 (E12, C6, black squares), T133 (G11, B4, red squares) and T145 (A8, C8, blue squares). Given the C<sub>1</sub> of each sample, the initial copy number is inferred from the standard curve obtained during the same experiment. Triplicate plots were performed for each tumor sample, but the data for only one are shown here. The results are shown in Table II.

30-fold maximum) (Berns et al., 1992; Borg et al., 1992; Courjal et al., 1997). (v) The erbB2 copy numbers obtained with real-time PCR were in good agreement with data obtained with other quantitative PCR-based assays in terms of the frequency and degree of amplification (An et al., 1995; Deng et al., 1996; Valeron

et al., 1996). Our results also correlate well with those recently published by Gelmini et al. (1997), who used the TaqMan system to measure erbB2 amplification in a small series of breast tumors (n = 25), but with an instrument (LS-50B luminescence spectrometer, Perkin-Elmer Applied Biosystems) which only allows end-

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Table II – Examples of cend! Gene dosage results from 3 breast tumors!

	eendl			aib			
Tumor	Capy	Mean	OZ	Cupy	Mcan	SD	Ncend I /alb
T118	4525			4223			
	4605	4603	77	4365	4325	89	1.06
	4678			4387			
T133	59821			9787			
	61659	61100	1111	10092	10137	375	6.03
	61821			10533			
T145	128563			7321			
	125892	125392	3448	7762	7672	316	16.34
	121722			7933			

<sup>1</sup>For each sample, I replicate experiments were performed and the mean and the standard deviation (SD) was determined. The level of cend1 gene amplification (Necnd1/alb) is determined by dividing the average cend1 copy number value by the average alb copy number value.

point measurement of fluorescence intensity. Here we report myc and cend! gene dosage in breast cancer by means of quantitative PCR. (vi) We found a high degree of concordance between real-time quantitative PCR and Southern blot analysis in terms of gene amplification, especially for samples with high copy numbers ( $\geq$ 5-fold). The slightly higher frequency of gene amplification (especially cend! and erbB2) observed by means of real-time quantitative PCR as compared with Southern-blot analysis may be explained by the higher sensitivity of the former method. However, we cannot rule out the possibility that some tumors with a few extra

gene copies observed in real-time PCR had additional copies of an arm or a whole chromosome (trisomy, tetrasomy or polysomy) rather than true gene amplification. These 2 types of genetic alteration (polysomy and gene amplification) could be easily distinguished in the future by using an additional probe located on the same chromosome arm, but some distance from the target gene. It is noteworthy that high sene copy numbers have the greatest prognostic significance in breast carcinoma (Borg et al., 1992; Slamon et al., 1987).

Finally, this technique can be applied to the detection of gene deletion as well as gene amplification. Indeed, we found a decreased copy number of erbB2 (but not of the other 2 protoncogenes) in several tumors; erbB2 is located in a chromosome region (17q21) reported to contain both deletions and amplifications in breast cancer (Bieche and Lidereau, 1995).

In conclusion, gene amplification in various cancers can be used as a marker of pre-neoplasia, also for early diagnosis of cancer, staging, prognostication and choice of treatment. Southern blotting is not sufficiently sensitive, and FISH is lengthy and complex. Real-time quantitative PCR overcomes both these limitations, and is a sensitive and accurate method of analyzing large numbers of samples in a short time. It should find a place in routine clinical gene dosage.

#### **ACKNOWLEDGEMENTS**

RL is a research director at the Institut National de la Santé et de la Recherche Médicale (INSERM). We thank the staff of the Centre René Huguenin for assistance in specimen collection and patient care.

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